JP,4101632,B [CLAIMS]

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CLAIMS

(57)[Claim(s)]

A polyoxyalkylene series polymer (A) which has a silicon containing functional group which can

construct a bridge by forming a siloxane bond,

Silicate (B)

an amine compound — and

As a silanol condensation catalyst, they are carboxylic acid tin salt (C) and a non-tin catalyst (E).

is a hardenability constituent to contain,

Silicate (B) is tetra alkoxysilane or its partial hydrolysis condensate,

A carbon atom in which <u>carboxylic acid</u> tin salt (C) adjoins a carbonyl group is the carboxylic acid tin

salt which is the 4th class carbon,

A carbon atom in which a non-tin catalyst (E) adjoins a carbonyl group is carboxylic acid which is the 4th class carbon.

A hardenability constituent.

[Claim 2]

A polyoxyalkylene series polymer in which a polyoxyalkylene series polymer (A) which has a silicon containing functional group which can construct a bridge by forming a siloxane bond introduced an unsaturation group into an end, and a general formula (1):

 $H-(SiR^{1}_{2-b}X_{b}O)_{m}-SiR^{2}_{3-a}X_{a}(1)$

the hardenability constituent acco<u>rding to claim 1 being</u> a polyoxyalkylene series polymer obtained by an addition reaction with a hydrosilane compound expressed. hydrolytic basis, and when two or more X exists, they may be the same and may differ. As for a, 0, 1, 2, or 3b shows 0, 1, or 2, respectively. About $\mathfrak b$ in $\mathfrak m$ bases (SiR 1 $_2$ - $_b$ X $_b$ O), they may be the same and may differ. m shows an integer of 0 to 19, however — what satisfies a+sigma b>=1 — carrying out different. When the Tori ORGANO siloxy group shown by aryl group of the carbon numbers 6-20, an an alkyl group of the carbon numbers 1-20 from which R¹ in a formula and R² were the same as or numbers 1-20 here, and three R may be the same and may differ. X shows a hydroxyl group or a aralkyl group of the carbon numbers 7–20, or {R?) $_3$ SiO– is shown and ${\sf R}^1$ or two or more ${\sf R}^2$ exist, they may be the same and may differ. R' is a hydrocarbon group of monovalence of the carbon

The hardenability constituent according to claim 1 or 2 in which a polyoxyalkylene series polymer (A) which has a silicon containing functional group which can construct a bridge by forming a siloxane bond is characterized by being a polymer which does not contain an amide segment (-NH-CO-) substantially in a principal chain skeleton.

An object for interior panels containing a hardenability constituent of a description in any 1 paragraph of Claims 1-3, an object for face panels, or adhesives for car panels. A sealing material for working joint of a building containing a hardenability constituent of a description in any 1 paragraph of Claims 1-3. http://www4.ipdl.inpit.go.jp/cgi-bin/tran_web_cgi_ejje?atw_u=http://www4.ipdl.inpit.go.jp... 2010/04/30

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DETAILED DESCRIPTION

[Detailed Description of the Invention]

Field of the Invention]

This invention relates to the hardenability constituent containing the organic polymer which has a silicon containing functional group (henceforth a reactive silicon group) which can construct a bridge by forming a siloxane bond.

Description of the Prior Art

It is known that the organic polymer which contains at least one reactive silicon group in a molecule accompanied by the hydrolysis reaction of a reactive silicon group, etc., and a rubber-like hardened has the interesting character in which construct a bridge by formation of the siloxane bond material is obtained with hygroscopic surface moisture etc. also in a room temperature.

polyisobutylene system polymer are already produced industrially, and are widely used for uses, such In the polymer which has these reactive silicon groups, a polyoxyalkylene series polymer and a as a sealing material, adhesives, and a paint.

nay pass with prudence and the stress from the outside of adherend, it may change by the time, and wall, the adhesives for car panels, etc. is inferior to stability or creep resistance, an adhesives layer inferior to stability or creep resistance, an adhesives layer may pass and it may change by the time, a panel tile, a stone, etc. may shift. Also in ceiling finishing adhesives or floor finishing adhesives, if electric equipment, an electron, and the adhesives for precision-mechanical-equipment assemblies adhesives for stone tensions, When the resin for adhesives used for the adhesives for finishing of and unevenness of a ceiling surface or a floor line may arise. If the stability of the electrical and The adhesives for interior panels, the adhesives for face panels, the adhesives for tiling, the

A sealing material generally fills up the joined part and crevice between various members, and he is these adhesives is excellent in stability or creep resistance.

and creep resistance are bad, an adhesives layer may pass, and it may change by the time, and may

oe connected with the degradation of apparatus. Therefore, it is called for that the constituent for

over a long period of time is very important, excelling in stability or endurance is called for as physical for a sealing material, the sealing material for direct grazing, the sealing material for multiple glass, the wall, and various face panels — business --- stability and endurance excellent in the constituent used width (Kasagi) the circumference of glass, the circumference of a window frame and a sash, a curtain used in order to give watertight and airtightness. Therefore, since the flattery nature to the use part properties of a hardened material. Working joint of a building with an especially large change of joint sealing material for speed signal generator construction methods, etc. are oalled for.

documents 7), the (patent documents 8), (The patent documents 9), the (patent documents 10), the (patent documents 11), the (patent documents 12), (The patent documents 13), the (patent documents 14), the (patent documents 15), the (patent documents 17), On the other hand, (the patent documents 1), the (patent documents 2), the (patent documents 3), the (patent documents 4), (The patent documents 5), the (patent documents 6), the (patent

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JP,4101632,B [DETAILED DESCRIPTION]

mainly indicated, and the description which suggests stability, creep resistance, and endurance is not constituent which uses as an essential ingredient the organic polymer which has the reactive silicon group which three hydrolytic bases combined on silicon is indicated, In these advanced technology, documents 21), the (patent documents 22), the (patent documents 23), the (patent documents 24) the (patent documents 18), the (patent documents 19), the (patent documents 20), In (the patent the fast curability based on the reactive silicon group which three hydrolytic bases combined is the (patent documents 25), the (patent documents 26), the (patent documents 27), the (patent documents 28), and the (patent documents 29), Although the room-temperature-curing nature indicated.

Patent documents 1]

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JP,H10-245482,A

Patent documents 2] [8000]

JP,H10-245484,A 6000

[Patent documents 3] JP,H10-251552,A

[Patent documents 4] [0010]

JP,H10-324793,A

Patent documents 5] [0011]

JP,H10-330630,A [0012]

Patent documents 6] JP,H11-12473,A

[0013]

Patent documents 7]

JP,H11-12480,A [0014]

Patent documents 8] JP,H11-21463,A

[0015]

[Patent documents 9] JP,H11-29713,A

Patent documents 10] 0016

JP,H11-49969,A

[Patent documents 11] [0017]

JP,H11-49970,A

Patent documents 12] JP,H11-116831,A [0018]

[0019]

Patent documents 13]

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Patent documents 14] NO No. 47939 [98 to]

[0021]

[Patent documents 15] JP,2000-34391,A

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[Patent documents 16] JP,2000-109676,A

Patent documents 17]

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Patent documents 20]

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Patent documents 27] JP,2001-72854,A 0034

Patent documents 28]

JP,2001-72855,A

Patent documents 29]

JP,2000-32771,A

Problem(s) to be Solved by the Invention

An object in view of the above-mentioned actual condition of this invention is to provide the stability, beiling finishing adhesives, Floor finishing adhesives, the adhesives for finishing of wall, the adhesives nterior panels with which stability, endurance, and creep resistance have been improved as for this invention, The adhesives for face panels, the adhesives for tiling, the adhesives for stone tensions, for car panels, the electrical and electric equipment, an electron and the adhesives for precisionmechanical—equipment assemblies, It aims at providing the sealing material for direct grazing, the provide the hardenability constituent which can give the hardened material excellent in stability, endurance, and creep resistance corrective strategy of a hardened material. The adhesives for methods, or the sealing material for working joint of a building. An object of this invention is to sealing material for multiple glass, the sealing material for speed signal generator construction endurance, and creep resistance. http://www4.ipdl.inpit.go.jp/cgi-bin/tran_web_cgi_ejje?atw_u=http%3A%2F%2Fwww4.ipdl.i... 2010/04/30

JP,4101632,B [DETAILED DESCRIPTION]

[Means for Solving the Problem]

By using on silicon a silicon containing functional group which has three or more hydrolytic bases as invention persons may solve such a problem, It found out improving stability, endurance, and creep a reactive silicon group of this polymer, as a result of inquiring wholeheartedly, in order that this resistance, and this invention was completed.

hardened material using an organic polymer (A) which has a silicon containing functional group which can construct a bridge by forming a siloxane bond of this invention, and a hardenability constituent That is, the 1st is related with stability, endurance, and creep resistance corrective strategy of a containing silicate (B).

The 2nd is an organic polymer which has a silicon containing functional group which can construct a containing an organic polymer (A1) which is a silicon containing functional group which has three or bridge by forming a siloxane bond of this invention, A silicon containing functional group which can resistance corrective strategy of a hardened material using on silicon a hardenability constituent construct a bridge by forming a siloxane bond, It is related with stability, endurance, and creep more hydrolytic bases,

endurance, and creep resistance corrective strategy of a hardened material given in the above using construct a bridge by forming a siloxane bond as a desirable embodiment, It is related with stability, A main chain of an organic polymer (A1) which has a silicon containing functional group which can a hardenability constituent which is an acrylic ester system copolymer manufactured by a living radical-polymerization method (meta).

It is related with stability, endurance, and creep resistance corrective strategy of a hardened material given in said either using a hardenability constituent which contains silicate (B) further as a desirable embodiment.

lt is related with stability, endurance, and creep resistance corrective strategy of a hardened material given in said either using a hardenability constituent which contains carboxylic acid tin salt (C) further as a desirable embodiment.

It is related with stability, endurance, and creep resistance corrective strategy of a hardened material given in said either using a hardenability constituent which contains an organic tin catalyst (D) further as a desirable embodiment.

The 3rd is an organic polymer which has a silicon containing functional group which can construct a functional group which can construct a bridge by forming a siloxane bond, and has it 1.7-5 pieces. endurance, and creep resistance corrective strategy of a hardened material using a hardenability bridge by forming a siloxane bond of this invention, This organio polymer is related with stability, constituent which is an organic polymer (A2) which averages per molecule a silicon containing

desirable embodiment, It is related with stability, endurance, and creep resistance corrective strategy A sílicon containing functional group which can construct a bridge by forming a siloxane bond as a of a hardened material given in the above using on silicon a hardenability constituent which is a silicon containing functional group which has three or more hydrolytic bases.

An organic polymer in which an organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond as a desirable embodiment introduced an unsaturation group into an end, and general formula (1):

H-(SiR¹ _{2-b}X_bO) _m-SiR² _{3-a}X_a (1)

an alkyl group of the carbon numbers 1–20 from which $\mathbb{R}^{\text{lin formul} a}$ and \mathbb{R}^2 were the same as or different. When the Tori ORGANO siloxy group shown by aryl group of the carbon numbers 6–20, an aralkyl group of the carbon numbers 7–20, or (R') $_3$ SiO– is shown and R 1 or two or more R 2 exist, 3/64 ふーツ

nay differ. m shows an integer of 0 to 19, however -- what satisfies a+sigma b>=1 -- carrying out -nydrolytic basis, and when two or more X exists, they may be the same and may differ. As for a, 0, 1, 2, or 3b shows 0, 1, or 2, respectively. About b in m bases (SiR 1 _{2-b}X $_b$ O), they may be the same and

it is related with stability, endurance, and creep resistance corrective strategy of a hardened material given in said either using a hardenability constituent which is an organic polymer obtained by an addition reaction with a hydrosilane compound expressed.

An organic polymer in which an organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond as a desirable embodiment introduced an unsaturation group into an end, and general formula (2): +-SiX, (2)

[0047]

hardened material given in said either using a hardenability constituent which is an organic polymer (X in a formula shows a hydroxyl group or a hydrolytic basis, and three X may be the same and) may differ. It is related with stability, endurance, and creep resistance corrective strategy of a obtained by an addition reaction with a hydrosilane compound expressed.

oridge by forming a siloxane bond of this invention, and this organic polymer is a general formula (3).: The 4th is an organic polymer which has a silicon containing functional group which can construct a -0-R3-CH(CH3)-CH2-(SiR1 2-bXb0) m-SiR2 3-aXa (3)

chosen from a group which consists of hydrogen, oxygen, and nitrogen as a composition atom) [show and] \mathbb{R}^1 , \mathbb{R}^2 , X, a, b, and m — the above — it is the same — it is related with stability, endurance, R^{3in formula} a divalent organic group of the carbon numbers 1-20 which contain one or more sorts and oreep resistance corrective strategy of a hardened material using a hardenability constituent which is an organic polymer (A3) which has a structure part with which it is expressed.

An organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond as a desirable embodiment is a general formula (4). -0-R3-C(CH3) =CH2 (4)

An organic polymer which introduced an unsaturation group expressed with $(\mathbb{R}^3$ is the same as the above), and general formula (1):

H-(SiR¹,

it is related with stability, endurance, and creep resistance corrective strategy of a hardened material addition reaction with a hydrosilane compound expressed with (R^{1 in formula} , R², X, a, b, and m are the given in the above using a hardenability constituent which is an organic polymer obtained by an _{2-b}X_bO) _m-SiR² _{3-a}X_a (1)

An organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond as a desirable embodiment is a general formula (5). [0020]

-O-R3-CH(CH3)-CH2-SIX3 (5)

same as the above).

(R^{3in formula} and X are the same as the above.) — it is related with stability, endurance, and creep resistance corrective strategy of a hardened material given in said either using a hardenability constituent which is an organic polymer which has a structure part with which it is expressed.

forming a siloxane bond as a desirable embodiment, It is related with stability, endurance, and creep An organic polymer which has a silicon containing functional group which can construct a bridge by resistance corrective strategy of a hardened material given in said either using for a principal chain skeleton a hardenability constituent which is an organic polymer which does not contain an amide segment (-NH-CO-) substantially

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A silicon containing functional group which can construct a bridge by forming a siloxane bond as a desirable embodiment is a general formula (6). :

- Si(OR4) 3 (6)

formula, respectively.) — it is related with stability, endurance, and creep resistance corrective strategy of a hardened material given in said either using a hardenability constituent which is a basis (three ${\sf R}^4$ is an organic group of monovalence of the carbon numbers 2-20 independently among a expressed.

forming a siloxane bond uses a hardenability constituent which is a triethoxy silyl group as a desirable It is related with stability, endurance, and creep resistance corrective strategy of a hardened material given in said either, wherein a silicon containing functional group which can construct a bridge by embodiment.

has three or more hydrolytic bases, and a hardenability constituent containing an organic tin catalyst strategy using on silicon an organic polymer (A1) which is a silicon containing functional group which The 5th is an organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond of this invention, A silicon containing functional group which can construct a bridge by forming a siloxane bond is related with thin layer hardenability corrective

An organic polymer (A) which has a silicon containing functional group which can construct a bridge precision-mechanical-equipment assemblies, a sealing material for direct grazing, a sealing material when the 6th forms a siloxane bond of this invention, And adhesives for interior panels containing for multiple glass, a sealing material for speed signal generator construction methods, or a sealing silicate (B), Adhesives for face panels, adhesives for tiling, adhesives for stone tensions, ceiling adhesives for car panels, the electrical and electric equipment, an electron and adhesives for finishing adhesives, It is related with floor finishing adhesives, adhesives for finishing of wall, material for working joint of a building.

sealing material for multiple glass, a sealing material for speed signal generator construction methods, The 7th is an organic polymer which has a silicon containing functional group which can construct a construct a bridge by forming a siloxane bond, . It is characterized by containing an organic polymer bridge by forming a siloxane bond of this invention, A silicon containing functional group which can stone tensions, Ceiling finishing adhesives, floor finishing adhesives, adhesives for finishing of wall, silicon. Adhesives for interior panels, adhesives for face panels, adhesives for tiling, adhesives for adhesives for car panels, It is related with the electrical and electric equipment, an electron and adhesives for precision-mechanical-equipment assemblies, a scaling material for direct grazing, (A1) which is a silicon containing functional group which has three or more hydrolytic bases on or a sealing material for working joint of a building.

mechanical-equipment assemblies, a sealing material for direct grazing, a sealing material for multiple finishing adhesives, floor finishing adhesives, adhesives for finishing of wall, adhesives for oar panels, polymerization method (meta) to the above by which it is characterized A description, Adhesives for interior panels, adhesives for face panels, adhesives for tiling, adhesives for stone tensions, Ceiling A main chain of an organic polymer (A1) which has a silicon containing functional group which can glass, a sealing material for speed signal generator construction methods, or a sealing material for construct a bridge by forming a siloxane bond as a desirable embodiment, Using a hardenability It is related with the electrical and electric equipment, an electron and adhesives for precisionconstituent which is an acrylic ester system copolymer manufactured by a living-radicalworking joint of a building. As a desirable embodiment, containing silicate (B) further to said either by which it is characterized A description, Adhesives for interior panels, adhesives for face panels, adhesives for stone tensions, Ceiling finishing adhesives, floor finishing adhesives, adhesives for finishing of wall, adhesives for car panels, It is related with the electrical and electric equipment, an electron and http://www4.ipdl.inpit.go.jp/cgi-bin/tran_web_cgi_ejje?atw_u=http%3A%2F%2Fwww4.ipdl.i.. 2010/04/30

adhesives for precision-mechanical-equipment assemblies, a sealing material for direct grazing, a sealing material for morthods, a sealing material for speed signal generator construction methods, or a sealing material for working joint of a building.

As a desirable embodiment, containing carboxylic acid tin salt (0) further to said either by which it is characterized A description, Adhesives for interior panels, adhesives for face panels, adhesives for tiling, adhesives for stone tensions, Ceiling finishing adhesives, floor finishing adhesives, adhesives for finishing of wall, adhesives for car panels, it is related with the electrical and electric equipment, an electron and adhesives for precision-mechanical-equipment assemblies, a sealing material for direct grazing, a sealing material for multiple glass, a sealing material for direct methods, or a sealing material for working joint of a building.

As a desirable embodiment, containing an organic tin catalyst (D) further to said either by which it is characterized A description, Adhesives for interior panels, adhesives for face panels, adhesives for tiling, adhesives for stone tensions, Ceiling finishing adhesives, floor finishing adhesives, adhesives for finishing of wall, adhesives for car panels, it is related with the electrical and electric equipment, an electron and adhesives for precision—mechanical—equipment assemblies, a sealing material for direct grazing, a sealing material for multiple glass, a sealing material for sortiuc onstruction methods, or a sealing material for working joint of a building.

The 8th is an organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond of this invention. It is characterized by this organic polymer being an organic polymer (A2) which averages per molecule a silicon containing functional group which can construct a bridge by forming a siloxane bond, and has it 1.7-5 pieces. Adhesives for interior panels, adhesives for face panels, adhesives for tiling, adhesives for stone tensions, Ceiling finishing adhesives, floor finishing adhesives, adhesives of mishing of wall, adhesives for car panels, it is related with the electrical and electric equipment, an electron and adhesives for precisionmechanical—equipment assemblies, a sealing material for direct grazing, a sealing material for multiple glass, a sealing material for speed signal generator construction methods, or a sealing material for precision methods, or a sealing material for precision methods.

A silicon containing functional group which can construct a bridge by forming a siloxane bond as a desirable embodiment. That it is a silicon containing functional group which has three or more hydrolytic bases on silicon to the above by which it is characterized A description, Adhesives for interior panels, adhesives for tiling, adhesives for stone tensions, Ceiling finishing adhesives, floor finishing adhesives, adhesives, adhesives for finishing of wall, adhesives for car panels, it is related with the electrical and electric equipment, an electron and adhesives for precision—mechanical-equipment assemblies, a sealing material for direct grazing, a sealing material for multiple glass, a sealing material for speed signal generator construction methods, or a sealing material for working joint of a building.

An organic polymer in which an organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond as a desirable embodiment introduced an unsaturation group into an end and a general formula (1) :

4-(SiR¹ _{2-b}X_bO) _m-SiR² _{3-a}X_a (1)

That it is an organic polymer obtained by an addition reaction with a hydrosilane compound expressed with (R^{1In} formula, R², X, a, b, and m are the same as the above) to said either by which it is characterized A description, Adhesives for interior panels, adhesives for face panels, adhesives for tiling, adhesives for stone tensions, Ceiling finishing adhesives, floor finishing adhesives, adhesives for finishing of wall, adhesives for car panels, it is related with the electrical and electric equipment, an electron and adhesives for precision—mechanical—equipment assemblies, a sealing material for multiple glass, a sealing material for multiple glass, a sealing material for working joint of a building.

סיסים. An organic polymer in which an organic polymer which has a silioon containing functional group which nttp://www4.ipdl.inpit_go.jp/cgi-bin/tran_web_cgi_ejje?atw_u=http\$3A%2F%2Fwww4.ipdl.i.. 2010/04/30

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oan oonstruct a bridge by forming a siloxane bond as a desirable embodiment introduced an unsaturation group into an end and a general formula (2) :

-SiX, (2)

That it is an organic polymer obtained by an addition reaction with a hydrosilane compound expressed with (X in a formula is the same as the above) to said either by which it is characterized A description, Adhesives for interior panels, adhesives for face panels, adhesives for tilling, adhesives for stone tensions, Ceiling finishing adhesives, floor finishing adhesives, for finishing of wall, adhesives for car panels, it is related with the electrical and electric equipment, an electron and adhesives for precision—mechanical—equipment assemblies, a sealing material for direct grazing, a sealing material for multiple glass, a sealing material for construction methods, or a sealing material for working joint of a building.

The 9th is an organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond of this invention, and this organic polymer is a general formula (3). : $-O-R^3-CH(CH_3)-CH_2-(SiR_{1-b}, V_bO)_m-SiR_{3-a}, V_a$ (3)

It is characterized by being an organic polymer (A3) which has a structure part expressed with (R lin formula, R², R³, X, a, b, and m are the same as the above). Adhesives for interior panels, adhesives for face panels, adhesives for tiling, adhesives for stone tensions, Ceiling finishing adhesives, floor finishing of wall, adhesives for car panels, It is related with the electrical and electric equipment, an electron and adhesives for precision—mechanical–equipment assemblies, a sealing material for direct grazing, a sealing material for multiple glass, a scaling material for speed signal generator construction methods, or a sealing material for working joint of a building. Inness

An organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond as a desirable embodiment is a general formula (4). :

 $-0-R^{1}-C(CH_{3})=CH_{2}$ (4)

An organic polymer which introduced an unsaturation group expressed with (R^1 is the same as the above), and general formula (1):

H-(SiR²_{2-b}X_bO) _m-SiR³_{3-a}X_a(1)

That it is an organic polymer obtained by an addition reaction with a hydrosilane compound expressed with (R^{2in formula}, R², X, a, b, and m are the same as the above) to the above by which it is characterized A description, Adhesives for interior panels, adhesives for face panels, adhesives for tiling, adhesives for stone tensions, Ceiling finishing adhesives, floor finishing adhesives, adhesives for finishing of wall, adhesives for oar panels, It is related with the electrical and electric equipment, an electron and adhesives for precision-mechanical-equipment assemblies, a sealing material for direct grazing, a sealing material for working joint of a building.

An organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond as a desirable embodiment is a general formula (5).

-0-R1-CH(CH3)-CH,-SiX3 (5)

(R^{1III} formula and X are the same as the above.) — a description to said either being an organic polymer which has a structure part with which it is expressed, Adhesives for interior panels, adhesives for face panels, adhesives for tiling, adhesives for stone tensions, Geiling finishing adhesives, floor finishing adhesives, adhesives for finishing of wall, adhesives for car panels, It is related with the electrical and electric equipment, an electron and adhesives for precision—mechanical—equipment assemblies, a sealing material for direct grazing, a sealing material for multiple glass, a sealing material for speed signal generator construction methods, or a sealing material for working joint of a building.

An organio polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond as a desirable embodiment. That it is an organic polymer which does not

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adhesives for tiling, adhesives for stone tensions, Ceiling finishing adhesives, floor finishing adhesives, adhesives for finishing of wall, adhesives for car panels, It is related with the electrical and electric naterial for direct grazing, a sealing material for multiple glass, a sealing material for speed signal contain an amide segment (-NH-CO-) substantially in a principal chain skeleton to said either by equipment, an electron and adhesives for precision-mechanical-equipment assemblies, a sealing which it is characterized A description, Adhesives for interior panels, adhesives for face panels, generator construction methods, or a sealing material for working joint of a building.

A silicon containing functional group which can construct a bridge by forming a siloxane bond as a desirable embodiment is a general formula (6). : [6900]

- Si(OR⁴) ₃ (6)

scaling material for multiple glass, a sealing material for speed signal generator construction methods, adhesives for precision-mechanical-equipment assemblies, a sealing material for direct grazing, a Adhesives for interior panels, adhesives for face panels, adhesives for tiling, adhesives for stone adhesives for car panels, It is related with the electrical and electric equipment, an electron and \mathbb{R}^{4in} formula is the same as the above.) -- a description to said either being a basis expressed, tensions, Ceiling finishing adhesives, floor finishing adhesives, adhesives for finishing of wall, a sealing material for working joint of a building.

description, Adhesives for interior panels, adhesives for face panels, adhesives for tiling, adhesives for sealing material for multiple glass, a sealing material for speed signal generator construction methods, desirable embodiment, That it is a triethoxy silyl group to said either by which it is characterized A A silicon containing functional group which can construct a bridge by forming a siloxane bond as a stone tensions, Ceiling finishing adhesives, floor finishing adhesives, adhesives for finishing of wall, adhesives for precision-mechanical-equipment assemblies, a sealing material for direct grazing, a adhesives for car panels, It is related with the electrical and electric equipment, an electron and or a sealing material for working joint of a building. The 10th is an organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond of this invention, A silicon containing functional group which can construct a bridge by forming a siloxane bond is related with an organic polymer (A1) which is a silicon containing functional group which has three or more hydrolytic bases on silicon, and a hardenability constituent containing silicate (B). [0072]

As a desirable embodiment, silicate is related with a hardenability constituent given in the above being a condensate of tetra alkoxysilane.

silicon containing functional group which has three or more hydrolytic bases on silicon, and a hardenability constituent containing carboxylic acid tin salt (C1) whose carbon of an alpha position of The 11th is an organic polymer which has a silicon containing functional group which can construct a construct a bridge by forming a siloxane bond, It is related with an organic polymer (A1) which is a bridge by forming a siloxane bond of this invention, A silicon containing functional group which can a carboxyl group is the 4th class carbon. The 12th is an organic polymer which has a silicon containing functional group which can construct a construct a bridge by forming a siloxane bond is related with an organic polymer (A1) and carboxylic bridge by forming a siloxane bond of this invention, A silicon containing functional group which can hydrolytic bases on silicon, and a hardenability constituent containing an organic tin catalyst (D). acid tin salt (G) which are the silioon containing functional groups which have three or more

The 13th is an organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond of this invention, A silicon containing functional group which can construct a bridge by forming a siloxane bond is related with an organic polymer (A1) which is a silicon containing functional group which has three or more hydrolytic bases on silicon, and a hardenability constituent containing a non-tin catalyst (E). http://www4.ipdl.inpit.go.jp/cgi-bin/tran_web_cgi_ejje?atw_u=http%3A%2F%2Fwww4.ipdl.i.. 2010/04/30

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he 14th is an organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond of this invention, A silicon containing functional group which can construct a bridge by forming a siloxane bond is related with an organic polymer (A1) which is a silicon containing functional group which has three or more hydrolytic bases on silicon, and a hardenability constituent containing a minute hollow body (F).

The 15th is an organic polymer which has a silicon containing functional group which can construct a on silicon, and is related with a hardenability constituent, wherein this organic polymer is 5 to 28 % of polymer (A1) which is a silicon containing functional group which has three or more hydrolytic bases construct a bridge by forming a siloxane bond, It is a hardenability constituent containing an organic bridge by forming a siloxane bond of this invention, A silicon containing functional group which can the weight in a total amount of a hardenability constituent. An organic polymer in which an organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond as a desirable embodiment introduced an unsaturation group into an end, and general formula (2):

It is related with a hardenability constituent given in said either being an organic polymer obtained by an addition reaction with a hydrosilane compound expressed with (X in a formula is the same as the above),

A silicon containing functional group which can construct a bridge by forming a siloxane bond as a desirable embodiment is a general formula (6):

- Si(OR⁴)₃ (6)

It is related with a hardenability constituent given in said either being a basis expressed with (R⁴ⁱⁿ formula is the same as the above).

containing functional group which can construct a bridge by forming a siloxane bond of this invention, and can construct a bridge by forming a siloxane bond is a general formula (6). : A silicon containing functional group which the 16th is an organic polymer which has a silicon

- Si(OR⁴) ₃ (6)

An organic polymer (A4) which has a basis expressed with (R⁴ⁱⁿ formula is the same as the above), and general formula (7)

- SiR⁵ (OR⁶) ₃₋₀ (7)

formula, respectively, and 3–c $\rm R^{6}$) It is an organic group of monovalence of the carbon numbers 2–20 which storage stability containing an aminosilane coupling agent (G) which has a basis expressed has independently, and c shows 0, 1, or 2, respectively. It is related with a hardenability constituent in (c ${
m R}^5$ is an organic group of monovalence of the carbon numbers 1-20 independently among a been improved.

A silicon containing functional group which the 17th is an organic polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond of this invention, and can construct a bridge by forming a siloxane bond is a general formula (6).:

- Si(OR⁴) ₃ (6)

An organic polymer (A4) which has a basis expressed with (R^{4In} form^{ula} is the same as the above), and general formula (8) :

- SiR7 (OCH3) (OR8) 3-d-e (8)

numbers 2-20 independently, respectively, d shows 0, 1, or 2 and e shows 1, 2, or 3.) However, 3-dformula, respectively, R⁸ of a 3-d-e individual is an organic group of monovalence of the carbon $(d\ R^7)$ is an organic group of monovalence of the carbon numbers 1–20 independently among a

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3>=0 shall be satisfied. It is a hardenability constituent containing an aminosilane coupling agent (H) which has a basis expressed, and is related with a hardenability constituent in which a cure rate ecuperating oneself beforehand in this hardenability constituent has been improved

containing functional group which can construct a bridge by forming a siloxane bond of this invention, A silicon containing functional group which the 18th is an organic polymer which has a silicon and can construct a bridge by forming a siloxane bond is a general formula (6).:

Si(OR4), (6)

it is related with an organic polymer (A4) which has a basis expressed with (R^{4in formula} is the same as the above), and a hardenability constituent containing epoxy resin (I). A silicon containing functional group which the 19th is a polyoxyalkylene series polymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond of this invention, and can construct a bridge by forming a siloxane bond is a general formula (6). :

Si(OR⁴) ₃ (6)

A polyoxyalkylene series polymer (A5) which has a basis expressed with (R4in formula is the same as the above), And it is related with a hardenability constituent containing an acrylic ester system copolymer (A6) which has a silicon containing functional group which can construct a bridge by forming a siloxane bond (meta).

which has a silicon containing functional group which can construct a bridge by forming a siloxane bond of this invertion, and can construct a bridge by forming a siloxane bond is a general formula (6). A silicon containing functional group which the 20th is a saturated hydrocarbon system polymer

- Si(OR4) 3 (6)

It is related with a hardenability constituent containing a saturated hydrocarbon system polymer (A7) which has a basis expressed with (R^{4in formula} is the same as the above). A silicon containing functional group which the 21st is an acrylic ester system copolymer which has a silicon containing functional group which can construct a bridge by forming a siloxane bond of this invention (meta), and can construct a bridge by forming a siloxane bond is a general formula (6). :

- Si(OR⁴)₃ (6)

lt is related with a hardenability constituent containing an acrylic ester system copolymer (A8) which nas a basis expressed with (R4in formula is the same as the above) (meta). [0086]

An organic polymer in which an organic polymer which has a silicon containing functional group which

can construct a bridge by forming a siloxane bond as a desirable embodiment introduced an

unsaturation group into an end, and general formula (9): H-Si(OR⁴)

It is related with a hardenability constituent given in said either being an organic polymer obtained by an addition reaction with a hydrosilane compound expressed with (being the same as the $\mathsf{R}^{4^{\mathsf{in}}}$ formula (6) 2 /

forming a siloxane bond as a desirable embodiment, It is related with a hardenability constituent given An organic polymer which has a silicon containing functional group which can construct a bridge by in said either being an organic polymer which does not contain an amide segment (-NH-CO-) [0087]

substantially in a principal chain skeleton.

A silicon containing functional group which can construct a bridge by forming a siloxane bond as a desirable embodiment is related with a hardenability constituent given in said either being a triethoxy

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containing functional group which can construct a bridge by forming a siloxane bond of this invention, A silicon containing functional group which the 22nd is an organic polymer which has a silicon and can construct a bridge by forming a siloxane bond is a general formula (6).

General formula (10) carrying out the ester exchange reaction of the compound (J) which has at least one methoxy group which can carry out an ester exchange reaction to an organic polymer (A4) which has a basis expressed with (R^{4in formula} is the same as the above) ;

- Si(OCH₃) _f(OR⁴) _{3-f} (10)

(among a formula, 3-f $\rm R^4$ is an organic group of monovalence of the carbon numbers 2-20 independently, respectively, and f shows 1, 2, or 3.) — it is related with a manufacturing method of an organic polymer which has a basis expressed

Hereafter, this invention is explained in detail

Restriction in particular does not have a principal chain skeleton of an organic polymer (A) which has a reactive silicon group used for this invention, and it can use a thing with various kinds of principal chain skeletons.

copolyamide, A diallyl phthalate system polymer etc. are illustrated. A polyoxyalkylene series polymer, polymerization of adipic acid and hexamethylenediamine, and sebacic acid, Nylon 11 by condensation carrying out condensation polymerization from polyamide system polymer,, for example, bisphenol A, butyl (meta) acrylate (meta); (meta) An acrylic ester system monomer, vinyl acetate, acrylonitrile, A polymerization of epsilon-aminoundecanoic acid, A polycarbonate system polymer manufactured by styrene; A vinyl monomer in inside of said organic polymer is polymerized. Graft polymer; obtained. Polysulfide system polymer:. Nylon 610 by condensation polymerization of nylon 6 by ring opening opening polymerization of epsilon-amino RAURO lactam, and the above-mentioned nylon, such as polymer obtained by ring opening polymerization of lactone, Ethyl (meta) acrylate, An acrylic ester vinyl-base polymer produced by carrying out the radical polymerization of the monomers, such as copolymer, a polycarbonate system polymer, etc. are preferred from acquisition and manufacture system copolymer produced by carrying out the radical polymerization of the monomers, such as polymers; Condensation with dibasic acid, such as adipic acid, and glycol, Or a polyester system polyisoprene, isoprene or butadiene, acrylonitrile, styrene, etc., A copolymer with polybutadiene, hydrogenation polyolefine system polymer produced by hydrogenating these polyolefine system and carbonyl chlorides which have a two or more-ingredient ingredient among Nylon 12 by ring połyoxypropylene polyoxy butylene copolymer, An ethylene-propylene system copolymer, A copolymer, A copolymer with copolymer of polyisobutylene, isobutylene, isoprene, etc., polychloroprene, A copolymer with a hydrocarbon system polymer, a polyester system polymer, an acrylic ester (meta) system Specifically A polyoxyethylene, polyoxypropylene, polyoxy butylene, Polyoxy tetramethylen, isoprene or butadiene, acrylonitrile, styrene, etc., Hydrocarbon system polymers, such as a polyoxyethylene polyoxypropylene copolymer, Polyoxyalkylene series polymers, such as a polymerization of epsilon caprolactam, hexamethylenediamine, Nylon 66 by condensation being easy among polymers with the above-mentioned principal chain skeleton.

system copolymer have a comparatively low glass transition temperature, and their hardened material Saturated hydrocarbon system polymers, such as polyisobutylene, hydrogenation polyisoprene, and hydrogenation polybutadiene, and a polyoxyalkylene series polymer and an acrylic ester (meta) obtained is preferred especially from excelling in cold resistance.

In a principal chain skeleton of the above-mentioned organic polymer (A), other ingredients, such as a urethane bond ingredient, may be included in the range which does not spoil an effect of this invention greatly.

is not limited especially as the above-mentioned urethane bond ingredient, but For example, toluene (tolylene) diisocyanate, Aromatic system polyisocyanates, such as diphenylmethane 2010/04/30 http://www4.ipdl.inpit.go.jp/cgi-bin/tran_web_cgi_ejje?atw_u=http%3A%2F%2Fwww4.ipdl.i...

disocyanate and xylylene disocyanate; Isophorone diisocyanate, What is obtained from a reaction of polyisocyanate compounds, such as aliphatic series system polyisocyanates, such as hexamethylene di-isocyanate, and polyol which has various kinds of above-mentioned principal chain skeletons can be mentioned.

[9600

If there are many amide segments (-NiH-CO-) generated in a principal chain skeleton based on said urethane bond, viscosity of an organic polymer will become high and will serve as a bad constituent of workability. Therefore, as for quantity of an amide segment occupied in a principal chain skeleton of an organic polymer, it is preferred that it is 3 or less % of the weight, it is more preferred that it is 1 or less % of the weight, and it is most preferred that an amide segment is not included substantially. [0097]

A reactive silicon group contained in an organic polymer which has a reactive silicon group is a basis which can construct a bridge by forming a siloxane bond by the reaction which has a hydroxyl group or a hydrolytic basis combined with a silicon atom, and is accelerated by a silanol condensation catalyst. As a reactive silicon group, it is a general formula (11).:

 $-(SiR^{1}_{2-b}X_{b}O)_{m}-SiR^{2}_{3-a}X_{a}(11)$

an alkyl group of the carbon numbers 1–20 from which R^{1in} formula and R^2 were the same as or different. When the Tori ORGANO siloxy group shown by aryl group of the carbon numbers 6–20, an aralkyl group of the carbon numbers 7–20, or $\langle R \rangle$ 3SiO- is shown and R^1 or two or more R^2 exist, they may be the same and may differ. R is a hydrocarbon group of monovalence of the carbon numbers 1–20 here, and three R may be the same and may differ. R shows a hydroxyl group or a hydrolytic basis, and when two or more X exists, they may be the same and may differ. As for a, 0, 1, 2, or 3b shows 0, 1, or 2, respectively. About b in m bases $\left\langle SiR^{\frac{1}{2}} z_{-b} X_b O\right\rangle$, they may be the same and may differ, m shows an integer of 0 to 19, however — what satisfies a+sigma b>=1 — carrying out — a basis expressed is raised.

LOUBRI
It is not limited but what is necessary is just a conventionally publicly known hydrolytic basis
Et is not limited but what is necessary is just a conventionan, a halogen atom, an alkoxy group, an
acyloxy group, a RETOKISHI mate group, an amino group, an amide group, an acid-amide group, an
aminooxy group, a sulfhydryl group, an alkenyloxy group, etc. are mentioned, for example. Among
these, a hydrogen atom, an alkoxy group, an acyloxy group, a KETOKISHI mate group, an amino group,
an amide group, an aminooxy group, a sulfhydryl group, and an alkenyloxy group are preferred,
hydrolysis nature is quiet and a viewpoint of handling or a cone to especially an alkoxy group is

preferred. [nnaq]

Louss).
A hydrolytic basis and a hydroxyl group can be combined with one silicon atom in the 1–3 ranges, and (a+sigmab) has 1–5 preferred ranges. When a hydrolytic basis and a hydroxyl group join together in [two or more] a reactive silicon group, they may be the same and may differ.

in particular, it is a general formula (12). :

0010

- SiR² 3-a X_a (12)

(R² and X are the same as the above among a formula.) Since a reactive silicon group expressed with an integer of 1–3 is easy to receive, a's is preferred.

[0101]

As an example of R¹ in the above-mentioned general formula (11) and (12), and R², For example, aralkyl groups, such as aryl groups, such as a methyl group, such as aryl groups, such as a methyl group and an ethyl group, and a cyclohexyl group, and a phenyl group, and benzyl, the Tori ORGANO siloxy group R is indicated to be by ₃SiO- which is a methyl group, a phenyl group, etc. (R), etc. are raised, Especially in those, a methyl group is preferred.

0102

As more concrete illustration of a reactive silicon group, a trimethoxysilyi group, a triethoxy silyl group, agroup, a dimethoxymethyl silyl group, and a group, and a

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diisopropoxy methyl silyl group are mentioned.

Especially in this invention, an organic polymer which has the silicon containing functional group (that is, the number of a+bxm of a general formula (11) is three or more) which three or more hydrolytic bases combined on silicon in an organic polymer of the (A) ingredient can be used as an ingredient

2

A hardened material which three or more hydrolytic bases had combined this (A1) ingredient on silicon, and constructed the bridge by a silanol condensation reaction of that reactive silicon group. Good stability is shown and remarkable creep resistance and an endurance improvement effect are shown as compared with a case of reactive silicon group containing organic polymer which has two or less hydrolytic bases.

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(A1) As for the number of a+bxm of a general formula (11) of an ingredient, it is more preferred that it is 3-5, and especially 3 is preferred. Also in it, since [that its improvement effect of the stability of a hardenability constituent of this invention, endurance, and oreep resistance is especially large and I the Tori alkoxy silyl groups has the good availability of a raw material, it is preferred. Thing of an alkoxyl group of the carbon numbers 1-20 is preferred, its thing of the carbon numbers 1-10 is more preferred, and its thing of the carbon numbers 1-4 is still more preferred here. Specifically, a trimethoxysilyl group and a triethoxy silyl group are the most preferred. Hardenability may become late when a carbon number is larger than 20.

Generally, if weight % of reactive silicon group containing organic polymer in a hardenability constituent becomes low, it is known that the endurance of a hardened material obtained will fall to **. However, if an ingredient (A1) of this invention is used as reactive silicon group containing organic polymer, high endurance is maintainable even if it makes low weight % of reactive silicon group containing organic polymer in a hardenability constituent. Therefore, five to 28% of the weight, when it is 15 to 24 % of the weight especially preferably, since a rate of an ingredient (A1) in a hardenability constituent is compatible in low cost and high endurance, it is more preferably preferred [rate] ten to 26% of the weight.

[0107]

processing in this invention, an organic polymer which has the Tori alkoxy silyl groups of the carbon numbers 2–20 can be used as a (A4) ingredient in an organic polymer of an ingredient (A1). Namely, general formula (6):

- Si(OR⁴) ₃ (6)

(three R^4 is an organic group of monovalence of the carbon numbers 2–20 independently among a formula, respectively.) — an organic polymer which has a basis expressed can be used as a (A4) ingredient.

<u>8</u>

It is known that methanol generated in connection with a hydrolysis reaction of a methoxy silyl group has peculiar toxicity of causing an obstacle of an optic nerve. On the other hand, since a carbon number of an alkoxy group which combines the (A4) ingredient with a silicon atom is 2 to 20, toxic high methanol is not contained in alcohol generated in connection with a hydrolysis reaction of a reactive silicon group, but serves as a constituent with high safety at it.

₽

(A4) It is preferred that it is especially 2-4, and when it is 2, it serves as ethanol, and since alcohol generated by hydrolysis has the highest safety, it is the most preferred [as for a carbon number of R⁴ of a general formula (6) of an ingredient, it is more preferred that it is 2-10, and / alcohol]. Specifically, a triethoxy silyl group is the most preferred. When a carbon number is larger than 20, while the hardenability of a hardenability constituent may become late, an anesthetic action and stimulation of alcohol to generate may be large.

[0110]

Especially in this invention, a principal chain skeleton can use as a (A5) ingredient what is polyoxyalkylene in an organic polymer of the (A4) ingredient. Namely, general formula (6):

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A polyoxyalkylene series polymer which has a basis expressed with $\langle \mathsf{R}^{4\mathsf{ln}} \, \mathsf{form}^{\mathsf{ula}} \, \mathsf{is}$ the same as the above) can be used as a (A5) ingredient.

effective network chain density of an organic polymer (A) ingredient contained in a hardened material 1.1-5 reactive silicon groups of an organic polymer (A) exist preferably [that average per molecule and at least one piece exists], and more preferably. If the number of reactive silicon groups silicon group may exist in an end of an organic polymer (A) chain, and may exist in an inside. Since nardened material in which a low elastic modulus is shown becomes is easy to be obtained by high ormed eventually will increase if a reactive silicon group exists in an end of a chain, a rubber-like become insufficient and will become difficult to reveal a good rubber elasticity action. A reactive contained in one molecule of organic polymers (A) will be less than one piece, hardenability will intensity and high elongation.

Especially in this invention, an organic polymer the number of reactive silicon groups per molecule averages, and 1.7–5 pieces exist in an organic polymer of the (A) ingredient can be used as an ingredient (A2).

reaction of that reactive silicon group, Good stability is shown, the number of reactive silicon groups per molecule averages, and remarkable creep resistance and an endurance improvement effect are A hardened material which the number of reactive silicon groups per molecule averaged this (A2) ngredient for it, and 1.7-5 pieces existed, and constructed the bridge by a silanol condensation shown as compared with a case of less than 1.7 organic polymers.

that they are 2-4 pieces, and it is preferred that they are especially 2.3-3 pieces. When there are few (A2) As for the number of reactive silicon groups per molecule of an ingredient, it is more preferred oonstituent of this invention, endurance, and creep resistance may not be enough, and when larger 1.7 reactive silicon groups per molecule, an improvement effect of the stability of a hardenability than five pieces, elongation of a hardened material obtained may become small.

Especially at this invention, it is a general formula in an organic polymer of the (A) ingredient (3). : -O-R3-CH(CH3)-CH2-(SIR12-6XbO) m-SIR23-4X4 (3)

chosen from a group which consists of hydrogen, oxygen, and nitrogen as a composition atom) [show R^{3In formula} a divalent organic group of the carbon numbers 1-20 which contain one or more sorts and] R¹, R², X, a, b, and m -- the above -- it is the same -- an organic polymer which has a structure part with which it is expressed can be used as a (A3) ingredient.

A hardened material which this (A3) ingredient has a structure part expressed with a general formula shows good stability, and shows remarkable creep resistance and an endurance improvement effect as compared with a case of an organic polymer which has terminal structures other than a general (3), and constructed the bridge by a silanol condensation reaction of that reactive silicon group

formula (3).

As for a carbon number of R³ of a general formula (3), it is more preferred from a point of availability that it is 1-10, and it is preferred that it is especially 1-4. Specifically, R³ has the most preferred

methylene group. [0118]

(A3) An ingredient is a general formula (5). :

-O-R3-CH(CH3)-CH2-SiX3 (5)

(R^{3in formula} and X are the same as the above.) — when it is an organic polymer which has a structure part with which it is expressed, since [that an improvement effect of the stability of a hardenability constituent of this invention, endurance, and creep resistance is especially large and] the availability

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of a raw material is good, it is desirable.

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(A) What is necessary is just to perform introduction of a reactive silicon group of an ingredient by a publicly known method. That is, the following methods are mentioned, for example.

group content epoxy compound. Subsequently, hydrosilane which has a reactive silicon group is made reactivity to an organic polymer which has functional groups, such as a hydroxyl group, in a molecule to this functional group react, and obtain an organic polymer containing an unsaturation group. Or an unsaturation group content organicity polymer is obtained by copolymerization with an unsaturation (b) Make an organic compound which has an active group and an unsaturation group which show to act on an acquired resultant, and it hydrosilylates.

(**) Make a compound which has a sulfhydryl group and a reactive silicon group react to an organic polymer containing an unsaturation group produced by making it be the same as that of the (b)

reactivity to an organic polymer which has functional groups, such as a hydroxyl group, an epoxy (**) Make a compound which has a functional group and a reactive silicon group which show group, and an isocyanate group, in a molecule to this functional group react.

Since a high inversion rate is obtained in comparatively short reaction time, a method of making a compound which has a polymer, an isocyanate group, and a reactive silicon group which have a hydroxyl group react to an end a method of (b) or among (***)s in the above method is preferred. An polymer obtained by a method and an organic polymer obtained by a method of (**) have the strong bad smell based on an mercaptosilane, especially its method of (b) is preferred. becoming a good hardenability constituent of workability by hypoviscosity rather than an organic organic polymer which has the reactive silicon group obtained by a method of (b), (**) Since

(b) As an example of a hydrosilane compound used in a method, For example, trichlorosilane, methyldi chlorosilicane, dimethylchlorosilicane, Halogenation Silang like phenyl dichlorosilane, Trimethoxysilane, dimethoxysilane; Methyldi acetoxysilane, The acyloxy silanes like a phenyldiacetoxysilane; although halogenation Silang and alkoxysilane are preferred, especially alkoxysilane has the quiet hydrolysis nature of a hardenability constituent obtained, and it is the most preferred to a handling and cone the KETOKISHI mate silanes like bis(dimethyl KETOKISHI mate)methylsilane and bis(cyclohexyl KETOKISHI mate)methylsilane are raised, it is not limited to these. Especially among these, Triethoxysilane, methyldiethoxysilane, methyl dimethoxysilane, The alkoxysilane like phenyl

In the above-mentioned hydrosilane compound, it is a general formula (2).

H-SiX₃ (2)

Since a hydrosilane compound expressed with (X in a formula is the same as the above) has an especially large improvement effect of the stability of a hardenability constituent which consists of an organio polymer obtained by an addition reaction of this hydrosilane compound, endurance, and creep trialkoxysilane, such as trimethoxysilane, triethoxysilane, and a triisopropoxy silane, is more preferred. resistance, it is preferred. In a hydrosilane compound expressed with a general formula (2),

group) of 1 also in said trialkoxysilane is like [when disproportionation may advance quickly and disproportionation progresses] dimethoxysilane — a dangerous compound arises in inside. From a viewpoint of safety on handling to a general formula (9) : trialkoxysilane in which carbon numbers, such as trimethoxysilane, have an alkoxy group (methoxy

H-Si(OR⁴)₃ (9)

It is preferred to use trialkoxysilane which has an alkoxy group whose carbon number expressed with $(\mathbb{R}^{4in}$ formula is the same as the above) is two or more. A viewpoint of availability, safety $\mathbb L$ on http://www4.ipdl.inpit.go.jp/ogi-bin/tran_web_ogi_ejje?atw_u=http%3A%2F%2Fwww4.ipdl.i.. 2010/04/30

handling], stability [of a hardenability constituent obtained], endurance, and creep resistance ** to triethoxysilane is the most preferred.
[0127]

(***) Although a method of introducing into an unsaturation binding site of an organic polymer a compound which has a sulfhydryl group and a reactive silicon group as a synthetic method by a radical addition reaction under a radical initiator and/or radical source-of-release existence, for example, etc. are mentioned, it is not limited in particular. As an example of a compound of having said sulfhydryl group and a reactive silicon group, for example, although gamma-mercapto propyltrimethoxysilane, gamma-mercaptopylmethyl dimethoxysilane, gamma-mercaptopropyl triethoxysilane, gamma-mercapto propylmethyl diethoxysilane, etc. are raised, it is not limited to these.

[0128]

(**) Although a method etc. which are shown in JP,H3–47825,A are mentioned, for example as a method of making a compound which has a polymer, an isooyanate group, and a reactive silicon group which have a hydroxyl group reacting to an end among synthetic methods, it is not limited in particular. As an example of a compound of having said isocyanate group and a reactive silicon group, For example, although gamma-isocyanate propyltrimethoxysilane, gamma-isocyanate propylmethyl dimethoxysilane, gamma-isocyanatepropyl triethoxysilane, gamma-isocyanate propylmethyl diethoxysilane, etc. are raised, It is not limited to these.

[0129]

As mentioned above, as for a silane compound which three hydrolytic bases have combined with one silicon atoms, such as trimethoxysilane, disproportionation may advance, As for trialkoxysilane in which especially carbon numbers, such as trimethoxysilane, have an alkoxy group (methoxy group) of 1, disproportionation may advance quickly. If disproportionation progresses, a remarkable dangerous compound [like] which is dimethoxysilane will arise. However, such disproportionation advances in neither gamma—mercapto propyltrimethoxysilane nor gamma—isocyanate propyltrimethoxysilane. For this reason, when using the Tori alkoxy silyl groups which has methoxy groups, such as a trimethoxysilyl group, as a silicon content group, it is preferred to use a synthetic method of (**) or (**).

[0130]

As how to obtain an organic polymer which has the silicon content group combined with a methoxy group, it is one method of the above-mentioned (**), (**), and (**), and a reactive silicon group is a general formula (6).

- Si(OR⁴)₃ (6)

A compound (J) which has at least one methoxy group which can carry out an ester exchange reaction after obtaining an organic polymer (getting it blocked the above-mentioned (A4) ingredient) which has a basis expressed with (R⁴in formula is the same as the above). It is a general formula by carrying out an ester exchange reaction under existence of a transesterification catalyst or nonexistence (10).:

- Si(OCH₃) _f(OR⁴) _{3-f} (10)

(among a formula, 3-f R⁴ is an organic group of monovalence of the carbon numbers 2-20 independently, respectively, and f shows 1, 2, or 3.) — a method of manufacturing an organic polymer which has a basis expressed can be mentioned. An organic polymer which has a basis expressed with a general formula (10) shows fast curability rather than an organic polymer which has a basis expressed with a general formula (6).

0131]

In said manufacturing method, after introducing a reactive silicon group especially by a method of (**), by carrying out an ester exchange reaction to the aforementioned (J) ingredient, A method of manufacturing an organic polymer which has a basis expressed with a general formula (10), Without a dangerous compound like dimethoxysilane by disproportionation arising in the middle of manufacture, there are few bad smells, and also since it becomes a good hardenability constituent of workability by hypoviscosity rather than an organic polymer obtained by a method of (***), it is more desirable than an organic polymer obtained by a method of (***).

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As a compound (J) which has at least one above-mentioned methoxy group which can carry out an ester exchange reaction, there is no limitation in particular and various kinds of compounds can be

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A compound etc. which have the silicon atom combined with methyl ester of various kinds of acid, such as methanol, carboxylio acid, and sulfonic acid, and at least one methoxy group as a (J) ingredient here can be mentioned. Since a compound which has the silicon atom united with 2-4 methoxy groups on the same silicon atom as a compound which has the silicon atom combined with said at least one methoxy group has a quick ester exchange reaction speed, it is preferred. Since especially a compound that has a silicon atom united with 2-4 methoxy groups on the same silicon atom and an amino group has a quick ester exchange reaction speed, it is preferred.

When it illustrates concretely, gamma-aminopropyl trimethoxysilane, gamma-aminopropyl methyl dimethoxysilane, gamma-(2-aminoethyl) aminopropyl trimethoxysilane, gamma-(2-aminoethyl) aminopropyl trimethoxysilane, gamma-ureido propyltrimethoxysilane, N-phenyl-gamma-aminopropyl trimethoxysilane, N-phenyl-gamma-aminopropyl trimethoxysilane, N-phenyl-gamma-aminopropyl trimethoxysilane, and n-benzyl-gamma-aminopropyl trimethoxysilane, can be mentioned. A denatured derivative and a condensation reaction thing of the above-mentioned silane compound as a (J) ingredient.

[0135]

Since an ester exchange reaction advances also under existence of a transesterification catalyst and comparatively low temperature conditions 60 ** or less, aforementioned amino group content Silang is preferred.

0126

As for the (J) ingredient used for this invertion, it is preferred to use and carry out an ester exchange reaction in 0.1–10 copies to 100 copies of reactive silicon group containing organic polymer of the (A4) ingredient. It is preferred to use it in 1–5 copies especially. The above–mentioned (J) ingredient may be used only by one kind, and may carry out two or more kind mixing use. [0137]

An organic polymer (A) which has a reactive silicon group may have straight chain shape or branching, and the number average molecular weight is 1,000–30,000 more preferably 500 to about 50,000 in polystyrene conversion in GPC. When there is a tendency whose number average molecular weight is inconvenient in respect of the extension characteristic of a hardened material at less than 500 and 50000 is exceeded, since it becomes hyperviscosity, there is an inconvenient tendency in respect of workability.

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A reactive silicon group may be in an end or an inside of an organio polymer ohain, and may be in both. Since effective network chain density of an organic polymer component contained in a hardened material formed eventually increases especially when a reactive silicon group is in a molecular terminal, it is desirable from points, like a rubber-like hardened material of high elongation becomes is easy to be obtained with high intensity.

[0139]

Said polyoxyalkylene series polymer is a general formula intrinsioally (13). : [0140]

[Formula 1]

 $-R^9-0-$ (16)

[0141]

(among a formula, R⁹ is a divalent organic group and is the straight chain shape or the branching alkylene group of the carbon numbers 1-14, — it being a polymer which has a repeating unit shown, and, R⁹ in a general formula (13) has the carbon numbers 1-14, and also preferred straight chain shape or branched state alkylene group of 2-4. As the example of the repeating unit shown by a general formula (13).

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Formula 2

** is mentioned. The principal chain skeleton of a polyoxyalkylene series polymer may consist of one kind of repeating unit, and may consist of two or more kinds of repeating units. When used especially for sealant etc., it is desirable from that what comprises the polymer which uses a propylene oxide polymer as the main ingredients is amorphous, or the point which is hypoviscosity comparatively. [0144]

tem, a U.S. Pat. No. 3278458 item, A U.S. Pat. No. 3278459 item, a U.S. Pat. No. 3427256 item, a U.S. As a synthetic method of a polyoxyalkylene series polymer, For example, the polymerizing method by shown in a U.S. Pat. No. 3427335 item etc., ********** such as the polymerizing method using a which are shown in JP,61-215623,A react, JP,46-27250,B, JP,59-15336,B, a U.S. Pat. No. 3278457 complex catalyst like a complex produced by making an organoaluminium compound and porphyrin Pat. No. 3427334 item, The polymerizing method by a composite metal oyanide complex catalyst polymerizing method using a catalyst which consists of a phosphazene compound illustrated by an alkali catalyst like KOH, the polymerizing method by a transition metal compound-porphyrin catalyst which consists of a polyphosphazene salt illustrated by JP,H10–273512,A, and the JP,H11-060722,A, are not limited in particular.

4366307, and U.S. Pat. No. 4960844, JP,61-197631,A, 61-215622, 61-215623, Although 6,000 or more JP,45-36319,B, 46-12154, JP,50-156599,A, 54-6096, 55-13767, 55-13468, What is proposed by each gazette, such as 57-164123, JP,3-2450,B, U.S. Pat. No. 3632557, U.S. Pat No. 4345053, U.S. Pat. No. JP,H3-72527,A, JP,H3-47825,A, and JP,H8-231707,A can illustrate a polyoxyalkylene series polymer with narrow molecular weight distribution in the 1.6 or less amount of Polymer Division, It is not number average molecular weights and Mw/Mn which are proposed by each gazette of 61–218632, A manufacturing method of a polyoxyalkylene series polymer which has a reactive silicon group, imited to in particular these.

A polyoxyalkylene series polymer which has the above-mentioned reactive silicon group may be used alone, and may be used together two or more sorts.

propylene, 1-butene, and isobutylene, is polymerized as a main monomer, and] (2) After making diene functional group into an end, and tend to control a molecular weight and can increase the number of isobutylene system polymer and a hydrogenation polybutadiene system polymer tend to introduce a series, such as butadiene and isoprene, homopolymerize or carrying out copolymerization of the end functional groups, they are preferred, and a composite ease to especially their isobutylene carbon-carbon unsaturated bonds other than an aromatic ring, and a polymer which makes the Said saturated hydrocarbon system polymer is a polymer which does not contain substantially above-mentioned olefinic compound, can obtain by a method of hydrogenating, but. Since an skeieton, (1). [whether an olefinic compound of the carbon numbers 1-6, such as ethylene, system polymer is preferred.

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JP,4101632,B [DETAILED DESCRIPTION]

which is excellent in heat resistance, weatherability, endurance, and humidity interception nature. That whose principal chain skeleton is a saturated hydrocarbon system polymer has the feature

All the monomeric units may be formed from an isobutylene unit, and an isobutylene system polymer,

originates in isobutylene from a field of rubber property 50% of the weight or more is preferred, what is contained 80% of the weight or more is more preferred, and especially a thing contained 90 to 99% Although a copolymer with other monomers may be sufficient, what contains a repeating unit which of the weight is preferred.

polymerized in 1.5 or less molecular weight distribution, and it is known that various functional groups As a synthetic method of a saturated hydrocarbon system polymer, although various polymerization methods are reported conventionally, especially living polymerization what is called of recent years isobutylene system polymer J. Polymer Sci, and Polymer Chem. Ed. 1997, By using 15 volumes and many is developed, an iniphor polymerization (J. -- P.Kennedy et al..) which was found out by 2843 pages, manufacturing easily is possible, about 500 to 100,000 molecular weight can be Kennedy and others in the case of a saturated hydrocarbon system polymer, especially an can be introduced into a molecular terminal.

example, JP,4-69659,B, JP,7-108928,B, JP,63-254149,A, Although it writes in each Description of JP,64-22904,A, JP,1-197509,A, Patent Gazette No. 2539445, Patent Gazette No. 2873395, and JP,7-As a process of a saturated hydrocarbon system polymer which has a reactive silicon group, For 53882,A, it is not limited to in particular these.

It is a general formula in a saturated hydrocarbon system polymer which has the above-mentioned reactive silicon group (6).:

- Si(OR4) 3 (6)

is the same as the above) can be used as an ingredient. This (A7) ingredient has the feature which is Especially (A7) a saturated hydrocarbon system polymer that has a basis expressed with (R⁴ⁱⁿ formula humidity interception nature of a principal chain skeleton, and does not have generation of methanol accompanying a hydrolysis reaction of a reactive silicon group, and is a polymer whose stability of a excellent in heat resistance based on a saturated hydrocarbon system polymer, weatherability, and hardened material, endurance, and creep resistance are still better.

A saturated hydrocarbon system polymer which has the above-mentioned reactive silicon group may be used alone, and may be used together two or more sorts.

Especially in this invention, a chain can use what is an acrylic ester (meta) system copolymer as an ingredient (A6) in an organic polymer of the (A) ingredient.

be used. If it illustrates, acrylic acid (meta), methyl acrylate (meta), (Meta) Ethyl acrylate, acrylic acid acid JIPA fluoromethylmethyl, acrylic acid (meta) 2-perfluoro methyl-2-perfluoro ethylmethyl, (Meta) (Meta) Acrylic acid-tert-butyl, acrylic acid (meta)-n-pentyl, (Meta) Acrylic acid-n-hexyl, acrylic acid acid (meta), (Meta) Acrylic acid trifluoromethyl methyl, an acrylic acid (meta) 2-trifluoro methylethyl, aforementioned (meta) acrylic ester system polymer, it is not limited but various kinds of things can butylethyl, (Meta) Acrylic acid 2-perfluoro ethyl, acrylic acid (meta) perfluoro methyl, (Meta) Acrylic Acrylic acid series (meta) monomers, such as acrylic acid 2-perfluoro hexylethyl, acrylic acid (meta) ethyl, (Meta) Acrylic acid-3-methoxy butyl, acrylic acid (meta)-2-hydroxyethyl, (Meta) Acrylic acidethylhexyl, acrylic acid (meta) nonyl, (Meta) Decyl acrylate, acrylic acid (meta) dodecyl, acrylic acid (meta)-n-propyl, acrylic acid (meta) isopropyl, (Meta) Acrylic acid-n-butyl, isobutyl acrylate (meta) (meta) cyclohexyl, (Meta) Acrylic acid-n-heptyl, acrylic acid (meta)-n-octyl, (Meta) Acrylic acid~2~ aminoethyl, gamma-(methacryloyl oxypropyl) trimethoxysilane, an ethyleneoxide addition of acrylic (meta) phenyl, (Meta) Acrylic acid toluyl, acrylic acid (meta) benzyl, acrylic acid (meta)-2-methoxy 2-hydroxypropyl, acrylic acid (meta) stearyl, metaglycidyl acrylate (meta), acrylic acid (meta) 2-Especially as an acrylic ester (meta) system monomer which constitutes a main chain of the (Meta) Acrylic acid 2-perfluoro ethylethyl, acrylic acid (meta) 2-perfluoro ethyl-2-perfluoro

such as methacrylamide; Alkenes; butadiene, such as vinyl ester; ethylene, such as vinyl acetate, vinyl ester monomer (meta) more preferably, is an acrylic polymer which consists of acrylic ester monomer aforementioned (meta) acrylic ester system copolymer, copolymerization of the following vinyl system preferred other monomers, copolymerization, and also that may carry out block copolymerization and preferably especially, and is a polymer which consists of butyl acrylates still more preferably. A point that physical properties, such as hypoviscosity of a compound, a low modulus of a hardened material, 2-perfluoro decylethyl, and acrylic acid (meta) 2-perfluoro hexadecylethyl, etc. are mentioned. In the ester and dialkyl ester of maleic acid and maleic acid; Fumaric acid, Monoalkyl ester and dialkyl ester acrylate is excellent in oil resistance. However, since it follows on increasing a ratio of butyl acrylate and the good oil resistance is spoiled, as for the ratio, for a use of which oil resistance is required, it monomers in that case. Acrylic acid (meta) expresses acrylic acid and/, or methacrylio acid with the monomers, such as KURORU styrene, styrene sulfonic acid, and its salt, Perfluoro ethylene, Fluoride are required is still more preferred. It can also transpose a part of ethyl acrylate to butyl acrylate in although limitation is not carried out, a copolymer of ethyl acrylate / butyl acrylate / acrylic acid 2preferred to use acrylic acid 2-methoxy ethyl, acrylic acid 2-ethoxyethyl, etc. by which oxygen was introduced into an alkyl group of a side chain. However, since it is in a tendency for heat resistance to be inferior by introduction of an alkoxy group which has an ether bond in a side chain, when heat resistance is required, it is preferred [the ratio] to make it to 40% or less. It is possible to obtain a Silicon content vinyl system monomers, such as vinyltriethoxysilane; A maleic anhydride, Monoalkyl preferred. It is an acrylic polymer which consists of acrylic ester monomer and a methacrylic-acid– Hexylmaleimide, octylmaleimide, dodecylmaleimide, stearyl maleimide, Maleimide system monomers, acrylate on the other hand in a use as which oil resistance, such as an automotive application, etc. such as phenylmaleimide and cyclohexylmaleimide; Acrylonitrile, Nitrile group content vinyl system order to improve the low-temperature characteristic etc., without spoiling oil resistance, it is also content vinyl monomers, such as perfluoro propylene and vinylidene fluoride; Vinyltrimetoxysilane, monomers, such as a methacrylonitrile; Acrylamide, Amide group content vinyl system monomers, is preferred to carry out to 40% or less, and also it is more preferred to make it to 30% or less. In order to raise that low-temperature characteristic, since it tends to be a little inferior to the lowtemperature characteristic (cold resistance), although a polymer mainly concerned with this ethyl polymer which changed the ratio and was suitable in consideration of physical properties needed, monomers can also be carried out with an acrylic ester (meta) system monomer. When this vinyl dienes, such as isoprene; VCM/PVC, a vinylidene chloride, an allyl chloride, allyl alcohol, etc. are mentioned. These may be used independently, and even if it carries out copolymerization of the propionate, vinyl pivalate, benzoic acid vinyl, and vinyl cinnamic acid, and propylene, Conjugated such as oil resistance, heat resistance, and the low-temperature characteristic, according to a various application or the purpose demanded. For example, as an example which is excellent in properties balance, such as oil resistance, heat resistance, the low-temperature characteristic, these desirable monomers are contained not less than 40% by a weight ratio in these desirable methoxy ethyl (it is 40-50/20-30/30-20 at a weight ratio) is mentioned. In this invention, it is of fumaric acid; Maleimide, Methylmaleimide, ethylmaleimide, propyl maleimide, butylmaleimide, monomer from physical properties etc. and (meta) an acrylic acid series monomer of output is plurality, they are not cared about. Especially, a polymer which consists of a styrene system system monomer is illustrated, styrene, vinyltoluene, alpha-methylstyrene, Styrene system above-mentioned expressive form.

acrylic ester system copolymer with low (meta) viscosity, and] to use a living-radical-polymerization distribution is narrow, and in order to obtain an acrylic ester system copolymer which has a crosscompound, a peroxide, etc. as a polymerization initiator generally has a value of molecular weight inking functional group in molecular chain terminals at a high rate (meta), it is preferred [it is an especially as a synthetic method of an acrylic ester system copolymer (A6). However, it has the distribution as large as two or more, and viscosity becomes high. Therefore, molecular weight (Meta) It is not limited but what is necessary is just to carry out by a publicly known method problem that a polymer obtained by the usual free radical polymerizing method using an azo

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above-mentioned "living-radioal-polymerization method" has halogen comparatively advantageous to halide compound An initiator, An "atom-transfer-radical-polymerization method" which polymerizes volumes, 5614 pages, etc. will be mentioned as this atom-transfer-radioal-polymerization method in an acrylic ester (meta) system monomer by making a transition metal complex into a catalyst, As a manufacturing method of an acrylic ester system copolymer which in addition to the feature of the Also in a "living-radical-polymerization method", an organic halogenated compound or a sulfonyl flexibility of a design of an initiator or a catalyst being large (meta), it is still more desirable. 117 a functional group conversion reaction etc. at the end, and has a specific functional group from Matyjaszewski et al. and journal OBU American chemical society (J. Am.Chem.Soc.) 1995.

As compared with an organic polymer which can improve notably the endurance of this (meta) acrylic transfer-radical-polymerization method", elongation may be insufficient and endurance may be bad. ester system copolymer by using on silicon a silicon containing functional group which has three or skeletons, such as a polyoxyalkylene series polymer. Even if it uses an acrylic ester (meta) system system copolymer which has a reactive silicon group (meta) may have low elongation as compared A hardened material produced by hardening a hardenability constituent containing an acrylic ester copolymer manufactured using the above "living-radical-polymerization method" and an "atomwith a hardenability constituent containing an organic polymer which has other principal chain more hydrolytic bases as a reactive silicon group, and has other principal chain skeletons, an endurance improvement effect is large.

process using the free radical polymerizing method for having used a chain transfer agent for JP,H3-14068,B, JP,H4-55444,B, JP,H6-211922,A, etc. is indicated, for example. Although a process which used an atom-transfer-radical-polymerization method for JP,H9-272714,A etc. is indicated, it is not As a process of an acrylic ester system copolymer which has a reactive silicon group (meta), a limited to in particular these.

It is a general formula in an acrylic ester system copolymer which has the above-mentioned reactive silicon group (meta) (6). :

- Si(OR⁴) ₃ (6)

accompanying a hydrolysis reaction of a reactive silicon group, and it is a polymer whose stability of a (meta) system copolymer of a principal chain skeleton in this (AB) ingredient, It has the feature which Especially (A8) an acrylic ester system copolymer that has a basis expressed with (R⁴in formula is the same as the above) (meta) can be used as an ingredient. Heat resistance based on an acrylic ester is excellent in weatherability and chemical resistance, and there is no generation of methanol hardened material, endurance, and creep resistance are still better.

component contained in a hardened material formed eventually increases especially when a reactive A reactive silioon group of the aforementioned (A8) ingredient may be in an end or an inside of an organic polymer chain, and may be in both. Since effective network chain density of a polymer silicon group is in an end of a polymer main chain, it is desirable from points, like a rubber-like hardened material of high elongation becomes is easy to be obtained with high intensity.

rate, it is more desirable and especially an atom-transfer-radical-polymerization method is preferred. polymerization method is used, since molecular weight distribution is narrow, it is hypoviscosity, and from the ability to introduce a cross-linking functional group into molecular chain terminals at a high As a polymerization method of the aforementioned (A8) ingredient, when a living-radicalAn acrylic ester system copolymer which has the above-mentioned reactive silicon group (meta) may be used alone, and may be used together two or more sorts.

together two or more sorts. Specifically, a polyoxyalkylene series polymer which has a reactive silicon An organic polymer which has these reactive silicon groups may be used alone, and may be used

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group, a saturated hydrocarbon system polymer which has a reactive silicon group, an acrylic ester system copolymer which has a reactive silicon group (meta), and an organic polymer which blends two or more sorts chosen from a group, ** and others, can also be used.

has a reactive silicon group, and an acrylic ester system copolymer which has a reactive silicon group (meta), Although proposed by JP,59-122541,A, JP,63-112642,A, JP,H6-172631,A, JP,H11-116763,A, A manufacturing method of an organic polymer which blends a polyoxyalkylene series polymer which etc., it is not limited to in particular these.

for an organic polymer which blends a polyoxyalkylene series polymer which has this reactive silicon group, and an acrylic ester system copolymer which has a reactive silicon group (meta) that stability it is known as compared with a case where a polyoxyalkylene series polymer is independently used is bad. Then, the above-mentioned general formula (6) as a polyoxyalkylene series polymer component in the aforementioned organic polymer to blend:

- Si(OR4) 3 (6)

has a reactive silicon group (meta) has outstanding stability, endurance, and creep resistance based A polyoxyalkylene series polymer (A5) which has a basis expressed with (R⁴ⁱⁿ form^{ula} is the same as the above) is used, An organic polymer blended with an acrylic ester system copolymer (A6) which on the (A5) ingredient, while outstanding weatherability and an adhesive property based on an ingredient (A6) are shown.

(A6) A desirable example of an acrylic ester (meta) system copolymer of an ingredient has a reactive silicon group, and a chain is a following general formula substantially (14).: [0167]

0168

(a hydrogen atom or a methyl group, and R^{11} show the alkyl group of the carbon numbers 1–8 among The acrylic ester monomer unit which has an alkyl group of the carbon numbers 1–8 expressed with a formula, as for $R^{10}\rangle$ (meta), and following general formula (15) :

Formula 4]

$$-CH_2-C-C-C-C-C-COR^{12}$$

[0171]

ten or more alkyl group ---> --- to a copolymer which consists of an acrylic ester monomer unit which (-- inside of formula, and R¹⁰ -- the above -- the same -- R¹² shows a with a carbon numbers of blending and manufacturing a polyoxyalkylene series polymer which has a reactive silicon group. has a with a carbon numbers of ten or more expressed alkyl group (meta-). It is the method of

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as R¹¹ of said general formula (14) --- the carbon numbers 1-8 of a methyl group, an ethyl group, a

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propy! group, n-butyl group, t-butyl group, a 2-ethylhexyl group, etc. --- desirable --- 1-4 --- an alkyl group of 1-2 is raised still more preferably. An alkyl group of R¹¹ may be independent and may be mixed two or more sorts.

as \mathbb{R}^{12} of said general formula (15) — ten or more carbon numbers of a lauryl group, a tridecyl group, a cetyl group, a stearyl group, a behenyl group, etc. -- usually --- 10-30 --- a long-chain alkyl group of 10–20 is raised preferably. Like a case of ${
m R}^{11}$, an alkyl group of ${
m R}^{12}$ may be independent and may be mixed two or more sorts.

monomeric unit of a formula (14) which exists in this copolymer, and a formula (15) surpasses 50 % of Although a chain of a ** (meta) acrylic ester system copolymer consists of a monomeric unit of a the weight. The sum total of a monomeric unit of a formula (14) and a formula (15) is 70 % of the formula (14) and a formula (15) substantially, a "real target" here means that the sum total of weight or more preferably.

As for an abundance ratio of a monomeric unit of a formula (14), and a monomeric unit of a formula (15), 95:5-40:60 are preferred at a weight ratio, and 90:10-60:40 are still more preferred.

ethyl methacrylate, A monomer containing amino groups, such as aminoethyl viryl ether; a monomerio unit which originates in acrylonitrile, styrene, alpha-methylstyrene, alkyl vinyl ether, VCM/PVC, vinyl groups, such as glycidyl acrylate and glycidyl methacrylate, Diethylamino ethyl acrylate, diethylamino formula (15), For example, acrylic acid, such as acrylic acid and methacrylic acid; Acrylamide, Amide As monomeric units other than a formula (14) which may be contained in this copolymer, and a groups, such as methacrylamide, N-methylolacrylamide, and N-methylolmethacrylamide, Epoxy acetate, vinyl propionate, ethylene, etc. in addition to this is raised.

has a reactive silicon group, and an acrylic ester system copolymer which has a reactive silicon group (meta) is proposed by JP,H1-168764,A, JP,2000-186176,A, etc., It is not limited to in particular these. Although an organic polymer which blends a saturated hydrocarbon system system polymer which

organio polymer which has a reactive silicon group as a manufacturing method of an organic polymer which blends an acrylic ester system copolymer which has a reactant silicon functional group (meta) A method of polymerizing an acrylic ester (meta) system monomer elsewhere under existence of an can be used. Although this manufacturing method is concretely indicated by each gazette, such as JP,59-78223,A, JP,59-168014,A, JP,60-228516,A, and JP,60-228517,A, it is not limited to these.

In this invention, silicate can be used as a (B) ingredient. This silicate has the function to improve the stability of an organic polymer which is the (A) ingredient of this invention, endurance, and oreep resistance.

[0180]

(B) Silicate which is an ingredient is a general formula (16).

Si(OR¹³) ₄ (16)

numbers 7–20 independently among a formula, respectively.) --- they are tetra alkoxysilane expressed $\langle \mathsf{R}^{13}$ are a hydrogen atom or an alkyl group of the carbon numbers 1–20, an aryl group of the carbon numbers 6-20, and the univalent hydrocarbon group chosen from an aralkyl group of the carbon or its partial hydrolysis condensate.

trimethoxysilane, dimethoxy diethoxysilane, methoxy triethoxysilane, Tetra alkoxysilane (tetraalkyl silicate), such as tetra n-propoxysilane, tetra i-propoxysilane, tetra n-butoxysilane, tetra ibutoxysilane, and tetra t-butoxysilane, and those partial hydrolysis condensates are raised. As an example of silicate, for example A tetramethoxy silane, a tetraethoxysilane, Ethoxy

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Since a partial hydrolysis condensate of tetra alkoxysilane has an improvement effect of the stability of this invention, endurance, and creep resistance larger than tetra alkoxysilane, it is preferred

A thing which was made to add and carry out partial hydrolysis of the water to tetra alkoxysilane by a usual method as a partial hydrolysis condensate of said tetra alkoxysilane for example, and was made ORGANO silicate compound. As such a condensate, the methylsilicate 51, the ethyl silicate 40 (all condense is raised. A commercial thing can be used for a partial hydrolysis condensate of an are made in Col Coat), etc. are mentioned, for example.

[0184]

Silicate (B) shows an improvement effect of still better stability, endurance, and creep resistance by combining with an ingredient (A1) of this invention, an ingredient (A2), and the (A3) ingredient. By combining with an ingredient (A1) especially shows an improvement effect of good stability, endurance, and oreep resistance.

range, an improvement effect of stability, endurance, and creep resistance may not be enough, and a cure rate may become slow if loadings of the (B) ingredient exceed this range. The above-mentioned section, and also 1 - 5 weight section is preferred. (B) If loadings of an ingredient are less than this (B) As amount of ingredient used, 0.1 - 10 weight section is preferred to (A) ingredient 100 weight silicate may be used only by one kind, and may carry out two or more kind mixing use.

resistance can be improved by using this carboxylic acid tin salt as a silanol condensation catalyst of silanol condensation catalysts, the stability of a hardened material obtained, endurance, and creep In this invention, carboxylic acid tin salt can be used as a (C) ingredient. As compared with other an organic polymer which is an ingredient (A1) of this invention. [0187]

Limitation in particular does not have carboxylic acid tin salt (C) used for this invention, and various kinds of compounds can be used for it.

including oarbonyl carbons, and carboxylic acid of a hydrocarbon system of the carbon numbers 2-20 As carboxylic acid which has an acid radical of carboxylic acid tin salt (C) here, a carboxylic acid group content compound of a hydrocarbon system of 2-40 is suitably used for a carbon number may be especially used suitably from a point of availability.

Enanthic acid, caprylic acid, 2-ethylhexanoic acid, pelargonic acid, capric acid, Undecanoic acid, lauric acid, 2-hydroxytetradecanoic acid, IPURORU acid, 2-hydroxyhexadecanoic acid, YARAPI Norian acid, 4.8,11,14-hexadeca tetraenoic acid, MOROKUCHI acid, steer RIDON acid, arachidonic acid, 8, 12 and unsaturated fatty acid, such as crotonic acid, isocrotonic acid, and 10-undecenoic acid; Reno elaidic acid, Linolic acid, 10,12-octadecadienoic acid, HIRAGO acid, alpha-eleostearic acid, beta-eleostearic and neo decanoio acid; PUROPI all acid, a tariric acid, Fatty acid with triple bonds, such as steer roll as gorlic acid; Acetoacetic acid, ethoxyacetic acid, Glyoxylic acid, glycolic acid, gluconic acid, sabinic Polyene unsaturated fatty acid, such as docosahexaenoic acid; 1-methylbutyric acid, Isobutyric acid, 2-ethylbutanoic acid, isovaleric acid, tuberculostearic acid, Branch fatty acid, such as a pivalic acid nalvalic acid, sterculic acid, HIDONO carbyne acid, chaulmoogric acid, Alicyclic carvone acids, such vaccenic acid, gadoleic acid, Gondo Inn acid, a cetoleic acid, erucic acid, brassidio acid, selacholeic acid, Tsuzuic acid, FIZETERIN acid, myristoleic acid, 2-hexadecenoic acid, 6-hexadecenoic acid, 7-When it illustrates concretely, acetic acid, propionic acid, butanoic acid, a valeric acid, caproio acid, acid, Straight chain saturated fatty acid groups, such as RAKUSERU acid; Undecylenic acid, Linder nonadecanoic acid, arachin acid, Behenic acid, lignoceric acid, cerinic acid, montanic acid, melissic hexadecenoic acid, palmitoleic acid, a petroselinic acid, Oleic acid, elaidic acid, ASUKUREPIN acid, 6, 19-docosatetraenoic acid, 4,8,12,15,18-eicosapentaenoic acid, clupanodonic acid, herring acid, acid, a crepenynic acid, KISHIMENIN acid, and 7-hexa crepe-de-Chine acid; Naphthenic acid, A punicic acid, linolenic acid, 8 and 11, 14-eicosatrienoic acid, A 7,10,13-docosatrienoic acid, acid, KISHIMEN acid, RUMEKUEN acid, acrylic acid, methacrylic acid, angelic acid, Monoene uni-PERIN acid, AMBURETTORU acid, ARYURITTO acid, 2-hydroxyoctadecanoic acid, 12acid, tridecylacid, myristic acid, pentadecyl acid, Pulmitic acid, heptadecylacid, stearic acid,

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itaconio aoid, etc. are mentioned. As aliphatic polycarboxylic acid, tricarboxylic acid, such as aconitic mentioned. In addition, amino acid, such as an alanine, leucine, threonine, aspartic acid, glutamic acid, isopropylbenzoic acid, salicylic acid, and toluic acid; aromatic polycarboxylic acids, such as phthalic chloroacrylic acid, and chlorobenzoic acid, etc. are mentioned. As aliphatic dicarboxylic acid, adipic acid, azelaic acid, pimelic acid, Saturation dicarboxylic acid, such as SUPERIN acid, sebacic acid, unsaturated dicarboxylic acid, such as maleic acid, fumaric acid, acetylene dicarboxylic acid, and anthracene carboxylic acid, Aromatic monocarboxylic acids, such as atrolactinic acid, anisic acid, acid, isophthalic acid, terephthalic acid, carboxyphenyl acetic acid, and pyromellitic acid, etc. are acid, citrate, and isocitric acid, etc. are mentioned. As aromatic carboxylic acid, benzoic acid, 9-Oxygenated fatty acid, such as recinoleic acid, cam ROREN acid, licanic acid, ferron acid, and cerebronic acid; a halogenation object of monocarboxylic acid, such as chloracetic acid, 2ethylmalonic acid, glutaric acid, oxalic acid, malonic acid, succinic acid, and oxydiacetic acid; hydroxyoctadecanoic acid, 18-hydroxyoctadecanoic acid, 9,10-dihydroxyoctadecanoic acid, arginine, cystein, methionine, phenylalanine, tryptophan, and histidine, is mentioned.

Especially acquisition is easy, and is cheap and a point that compatibility with an ingredient (A1) is good to said carboxylic acid has 2-ethylhexanoic acid, octylic acid, neo decanoic acid, oleic acid, or preferred naphthenic acid.

(workability --- bad). Therefore, as for the melting point of said carboxylic acid, it is preferred that it is when the melting point of said carboxylic acid is high (crystallinity is high), the melting point becomes 65 ** or less, it is more preferred that it is -50-50 **, and it is preferred that it is especially -40-35 high in a similar manner, and it is hard to deal with carboxylic acid tin salt which has the acid radical

liquefied and carboxylic acid tin salt which has the acid radical becomes a thing which has a high solid extended a constituent thinly, volatilization by heating may be large and catalyst ability of carboxylic acid metal salt may fall greatly. Therefore, as for said carboxylic acid, it is preferred that oarbon numbers including carbon of a carbonyl group are 2-20, it is more preferred that it is 6-17, and it is mostly an ingredient which volatilizes easily with heating. Especially on conditions (thin layer) which carboxylic acid metal salt may fall [carboxylic acid tin salt which has the acid radical], including state or viscosity and which is hard to deal with it (workability -- bad). On the contrary, when a when a carbon number of said carboxylic acid is large (a molecular weight is large), it becomes carbon number of said carboxylic acid is small (a molecular weight is small), catalyst ability of preferred that it is especially 8-12.

carboxylic acid tin salt to dicarboxylic acid or monocarboxylic acid, and it is more preferred that it is It is preferred that it is the tin salt of a point of the ease (workability, viscosity) of dealing with it of the tin salt of monocarboxylic acid.

As said monocarboxylic acid tin salt, it is a general formula (17). :

Sn(OCOR) 2 (17)

(The inside R of a formula is substitution or an unsubstituted hydrocarbon group, and may include a carbon carbon double bond.) Two RCOO-bases may be the same and may differ. A divalent Sn compound expressed or general formula (18): Sn(0COR) 4 (18)

(It is the same as the above the inside R of a formula.) Two RCOO-bases may be the same and may differ. A tetravalent Sn compound expressed is preferred. A divalent Sn compound expressed with a general formula (17) from a point of hardenability and availability is more preferred.

carboxylic acid tin salt (neo decanoic acid tin.) in which said carboxylic acid tin salt (C) is carboxylic acid tin salt (2—ethylhexanoic acid tin etc.) and the 4th class carbon whose carbon of an alpha position of a carboxyl group is the 3rd class carbon Pivalic-acid tin etc. are more preferred from a cure rate being quick, and especially carboxylic acid tin salt whose carbon atom which adjoins a carbonyl group is the 4th class carbon is preferred. http://www4.ipdl.inpit.go.jp/cgi-bin/tran_web_cgi_ejje?atw_u=http%3A%2F%2Fwww4.ipdl.i.. 2010/04/30

Especially in this invention, carbon of an alpha position of a carboxyl group uses carboxylic acid tin salt which is the 4th class carbon as an ingredient (C1) in carboxylic acid tin salt (C) [0197]

(C1) As carboxylic acid tin salt of an ingredient, it is a general formula (19).

[0198]

Formula 5

$$\begin{pmatrix}
R^{14} & Q \\
R^{15} - C - C - O \\
R^{16} & \end{pmatrix}_{2} (19)$$

[0199]

(among the formula, R¹⁴, R¹⁵, and R¹⁶ are the independent substitution or unsubstituted univalent organic groups, respectively, and may contain the carboxyl group.) -- the chain fatty acid tin expressed or general formula (20):

0500

[Formula 6]

0201

Among the formula, a substitution or unsubstituted univalent organic group and ${\sf R}^{18}$ are substitution or unsubstituted divalent organic groups, and ${\rm R}^{17}$ may contain the carboxyl group, respectively.) It reaches and is a general formula (21).

Formula 7] [0202]

[0203]

decanoic acid, BASA tic acid, Chain monocarboxylic acid, such as 2,2-dimethyl- 3-hydroxypropionic acid which has an acid radical of carboxylic acid tin salt (C1) is illustrated concretely, A pivalic acid, (among a formula, R¹⁹ is a substitution or unsubstituted trivalent organic group, and may contain a carboxyl group.) -- cyclic-fatty-acid tin containing structure expressed is mentioned. If carboxylic dimethylvaleric acid, a 2-ethyl-2-methylvaleric acid, a 2,2-diethylvaleric acid, 2,2-dimethylhexane scid, 2,2-dicthylhexanoic acid, 2,2-dimethyloctanoic acid, 2-ethyl-2,5-dimethylhexane acid, neo 2.2-dimethylbutanoio acid, 2-ethyl-2-methylbutyric acid, 2,2-diethylbutanoic acid, A 2,2http://www4.ipdl.inpit.go.jp/cgi-bin/tran_web_cgi_ejje?atw_u=http%3A%2F%2F%2Fwww4.ipdl.i.. 2010/04/30

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methyloyclohexane carboxylic acid, 2-methylbicyclo[2.2.1]-5- heptene-2-carboxylic acid, 2-methylacid, Dimethylmalonic acid, ethyl methylmalonic acid, diethylmalonic acid, 2,2–dimethyl amber acid, Chain dicarboxylic acid, such as 2,2–diethyl amber acid and 2,2–dimethylglutaric acid, Chain methyloyolopentanecarboxylic acid, 1,2,2-trimethyl 1,3-cyclopentane dicarboxylic acid, 1tricarboxylic acid, such as 3-methyliso citrate and 4.4-dimethylaconitic acid, 1-

etc. are mentioned. These can also be used aithough many compounds containing such a structure to 7-oxabicyclo [2.2.1]-5-heptene-2-carboxylic acid, 1-adamantane carboxylic acid, Annular carboxylic acid, such as bicyclo[2,2,1] heptane- 1-carboxylic acid and bicyclo[2,2,2] octane- 1-carboxylic acid, a natural product exist

acquisition is easy, pivalic-acid tin, neo decanoic acid tin, BASA tic acid tin, 2,2-dimethyloctanoic monocarboxylate is more preferred and also chain tin monocarboxylate is more preferred. Since From a point that compatibility with an ingredient and workability are especially (A1) good, tin acid tin, especially 2-ethyl-2,5-dimethylhexane acid tin, etc. are preferred.

(C1) Also in an ingredient, although carboxylate of divalent tin and carboxylate of tetravalent tin are mentioned like a case of the above-mentioned (C) ingredient, carboxylate of a point of hardenability and availability to divalent tin is more preferred.

preferred that it is 5–20, it is more preferred that it is 6–17, and it is preferred that it is especially 8– catalytic activity may fall if a carbon number increases more than this range, it is not desirable. It is 12. Since compatibility with an ingredient may fall that it is easy to become a solid state (A1) and As for a carbon number of carboxylic acid which has an acid radical of an ingredient (C1), it is not desirable from on the other hand, volatility, the increase of a smell, and the thin layer hardenability of a hardenability constituent falling, if there are few carbon numbers.

As an ingredient, from these points (C1) Neo decanoic acid tin (divalent), BASA tic acid tin (divalent), 2,2-dimethyloctanoic acid tin (divalent), 2-ethyl-2,5-dimethylhexane acid tin (divalent), Neo decanoic acid tin (tetravalence), BASA tic acid tin (tetravalence), 2,2-dimethyloctanoic acid tin (tetravalence), and especially 2-ethyl-2,5-dimethylhexane acid tin (tetravalence) are preferred.

(C) As amount of an ingredient and (C1) ingredient used, about 0.01–20 weight sections are preferred cure rate may become slow and a hardening reaction will become fully difficult to advance if loadings are less than this range, it is not desirable. On the other hand, if loadings exceed this range, working to ingredient (A1) 100 weight section, and also about 0.5-10 weight sections are preferred. Since a life becomes short too much, and workability may worsen, and it is not desirable from a point of storage stability.

The (C) ingredient and (C1) an ingredient can be used combining two or more sorts besides using it

On the other hand, only of the (G) ingredient and (G1) an ingredient, activity is low, and when

As various amine compounds, although indicated to JP,H5-287187,A, for example, Specifically moderate hardenability is not acquired, an amine compound can be added as a co-catalyst.

Triethylamine, Aliphatic series tertiary amines, such as triamylamine, trihexyl amine, and trioctylamine; ethylhexyl) amine, Didecyl amine, dilauryl amine, JISECHIRU amine, distearyl amine, Aliphatic series series primary amines, such as Sept Iles amine, stearylamine, and cyclohexylamine; Dimethylamine, octylamine, 2-ethylhexylamine, Nonyl amine, decyl amine, lauryl amine, pentadecyl amine. Aliphatic aniline, stearylaniline, a triphenylamine, N.N-dimethylaniline, dimethylbenzyl aniline, etc. reach, As Triaryi amine, Aliphatic series unsaturation amines, such as oleylamine; aromatic-amine; Lauryl Diethylamine, dipropyl amine, diisopropylamine, dibutyl amine, Diamylamine, dioctyl amine, di(2-Methylamine, ethylamine, propylamine, isopropylamine, A butylamine, amyl amine, hexylamine, secondary amines, such as methylstearylamine, ethylstearylamine, and butylstearylamine:

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other amines, monoethanolamine, diethanolamine, Triethanolamine, dimethylamino ethanol,

oadings of an amine compound exceed 20 weight sections, pot life may become short too much and weight sections, and a hardening reaction becomes fully difficult to advance. On the other hand, if preferred. A cure rate may become it slow that loadings of an amine compound are less than 0.01 As for loadings of said amine compound, about 0.01-20 weight sections are preferred to organic polymer 100 weight section of an ingredient (A1), and also its 0.1 to 5 weight section is more not preferred from a point of workability.

satalyst is used as a silanol condensation catalyst of an organic polymer which has a reactive silicon group, as compared with other silanol condensation catalysts, a hardenability constituent with high catalytic activity, and good depths hardenability and an adhesive property is obtained. However, in this invention, an organic tin catalyst can be used as a (D) ingredient. When this organic tin according to an addition of this organic tin catalyst, the stability of a hardened material of a nardenability constituent obtained, endurance, and creep resistance fall.

activity is high, and depths hardenability and an adhesive property are good, and the stability of a A hardenability constituent which added an organic tin catalyst of the (D) ingredient by using an organic polymer which is an ingredient (A1) of this invention as a polymer component, Catalytic hardened material obtained, endurance, and creep resistance can be maintained highly

combined, the hardenability of a thin layer part can be improved notably, maintaining the stability of a portion, and may remain on conditions of heat and high humidity especially with un-hardening. On the In using adhesives or a sealing material which, on the other hand, contains an organic polymer which other hand, if said organic tin catalyst (D) is used as a ouring catalyst, as mentioned above, stability catalyst of an organic polymer and the (D) ingredient which is an ingredient (A1) of this invention is nowever, if this carboxylic acid tin salt is used as a curing catalyst, when it will be alike around a masonry joint and a sealing material will remain by a thin layer, it is hard to harden that thin layer carboxylic acid tin salt of the aforementioned (C) ingredient as a curing catalyst in many cases. and endurance will fall, but the hardenability of a thin layer part is good. Then, if an organic tin has a reactive silicon group as the main ingredients for a use which needs endurance, it uses hardened material obtained, and endurance highly.

depending on an addition of an organio tin catalyst of the (D) ingredient, stability and endurance may (all a little. Then, it is more preferred to decrease the quantity of an addition of the (D) ingredient to However, even if it combines with an organic polymer which is an ingredient (A1) of this invention, hardenability, depths hardenability, an adhesive property, and thin layer hardenability are acquired such an extent that carboxylic acid tin salt of the (C) ingredient is used together and sufficient with an organic tin catalyst of the (D) ingredient as a curing catalyst. [0217]

As an example of said organic tin catalyst (D), they are dialkyl tin carboxylate, dialkyl tin oxide, and a general formula (22).

Q Sn(OZ) 4-g or [Q2Sn(OZ)] 20 (22)

expresses among a formula an organic group which has a functional group with which Q can form a diphenyldimethoxysilane, and phenyltrimethoxysilane. Since activity as a silanol condensation catalyst coordinate bond in an inside of a univalent hydrocarbon group of the carbon numbers 1–20, or self for a univalent hydrocarbon group of the carbon numbers 1-20 to Sn.) g is 0, 1, 2, or 3. A compound etc. diacetate, It is usable as a (D) ingredient also in a reactant with a low molecule silicon compound which are shown are shown. Tetravalent tin compounds, such as dialkyl tin oxide and dialkyl tin which has hydrolytic silicon groups, such as a tetraethoxysilane, methyl triethoxysilane,

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is high, chelate compound and tin alcoholates, such as a compound shown by a general formula (22)

also in these, i.e., dibutyl tin bisacetylacetonate etc., are more preferred

tin dibenzyl malate, dibutyltin maleate, dioctyl tin diacetate, dioctyl tin distearate, dioctyl tin dilaurate, As an example of said dialkyl tin carboxylate, For example, dibutyltin dilaurate, dibutyltin diacetate, a dibutyl tin diethylhexano rate, Dibutyl tin JIOKUTETO, dibutyl tin dinethylmalate, dibutyl tin diethyl malate, Dibutyl tin dibutyl maleate, dibutyl tin diisooctyl malate, Dibutyl tin ditridecyl malate, dibutyl

As an example of said dialkyl tin oxide, dibutyl tin oxide, dioctyl tin oxide, a mixture of dibutyl tin oxide dioctyl tin diethyl malate, dioctyl tin diisooctyl malate, etc. are mentioned

If said chetate compound is illustrated concretely,

and phthalic ester, etc. are mentioned.

Formula 8]

Although ** is mentioned, it is not limited to these. In these, its catalytic activity is high, and it is low cost, and since dibutyl tin bisacetylacetonate is easy to receive, it is the most preferred

If said tin alcoholates are illustrated concretely,

Formula 9]

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(C₄H₉) ₃Sn0CH₃

(C₄H₉) ₂Sn (0CH₃) ₂

C₄H₉Sn (0CH₃)₃

Sn (0CH₃) 4

(C₄H₉) ₂Sn (0C₃H₇) ₂

 $(C_4H_9)_2Sn(0C_4H_9)_2$

(G4H₉)₂Sn (0G₈H₁₇)₂

 (C_4H_9) ₂Sn $(0C_{12}H_{25})$ ₂

(C₈H₁₇)₂Sn (0CH₃)₂

 $(C_4H_9)_2Sn(0\langle)_2$

 $(C_4H_9)_2Sn(0)$

 $(C_4H_9)_2Sn(0(1))_2$

[(C4H₉)₂\$n]₂0

 $[(C_4H_9) _2\Sn]_20$

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Although ** is mentioned, it is not limited to these. In these, a dialkyl tin JIARUKOKI side is preferred. Especially the dibutyl tin JIMETOKI side is low cost, and since it is easy to receive, it is

range, it is not desirable. On the other hand, if loadings exceed this range, working life becomes short (D) As amount of ingredient used, about 0.01–20 weight sections are preferred to ingredient (A1) 100 weight section, and also about 0.1-10 weight sections are preferred. Since a cure rate may become slow and a hardening reaction will become fully difficult to advance if loadings are less than this too much, and workability may worsen, and it is not desirable from a point of storage stability.

exceed this range, working life becomes short too much and workability may worsen. (D) The stability When loadings of an ingredient are less than this range, a cure rate may become slow, when loadings ingredient 0.01 - 10 weight section to ingredient 100 weight section, and also it is more preferred to ingredient are less than this range, and loadings exceed this range, endurance, and creep resistance hardenability, an adhesive property, and thin layer hardenability may not be enough if loadings of an As amount of [in case used of using the (D) ingredient and the (C) ingredient together as a curing consider it as (C) ingredient 1-10 weight section and (D) ingredient 0.02-5 weight section. (G) of a hardened material which will be obtained if an improvement effect of hardenability, depths catalyst], (A1) It is preferred to consider it as (C) ingredient 0.5 - 20 weight section and (D)

The (D) ingredient can be used combining two or more sorts besides using it alone.

condensation catalyst of an organic polymer which is an ingredient (A1) of this invention. A non-tin In this invention, a non-tin catalyst can be used as a (E) ingredient. This non-tin catalyst has a function which improves the stability of a hardened material obtained, endurance, and creep catalyst which is the (E) ingredient is an eco-friendly curing catalyst with high social needs. resistance as compared with other silanol condensation catalysts, when it uses as a silanol

As a non-tin oatalyst which is the (E) ingredient which can be used for this invention, although there is no restriction in particular, an organio metallic compound containing carboxylic acid, carboxylic acid metal salt other than carboxylic acid tin salt, organic sulfonic acid, alkyl acid phosphate and 3B fellows, and 4A group metal, etc. are illustrated.

The various above-mentioned carboxylic acid which has an acid radical of carboxylic acid tin salt which is the (C) ingredient as carboxylic acid can be illustrated.

that it is especially 8–12. A point to dicarboxylic acid or monocarboxylic acid of the ease (workability, viscosity) of dealing with it of carboxylic acid is preferred, and monocarboxylic acid is more preferred including carbon of a carbonyl group are 2–20, it is more preferred that it is 6–17, and it is preferred carboxylic acid (neo decanoic acid.) in which said carboxylic acid is carboxylic acid (2-ethylhexanoic acid etc.) and the 4th class carbon whose carbon of an alpha position of a carboxyl group is the 3rd class carbon A pivalic acid etc. are more preferred from a cure rate being quick, and especially carboxylic acid whose carbon atom which adjoins a carbony! group is the 4th class carbon is As for said carboxylic acid, it is preferred like carboxylic acid tin salt (C) that carbon numbers preferred.

Especially as carboxylic acid, 2-ethylhexanoic acid, neo decanoic acid, BASA tic acid, 2,2-dimethyloctanoic acid, and 2-ethyl-2,5-dimethylhexane acid are preferred from a point of availability, hardenability, and workability. As carboxylic acid metal salt other than said carboxylic acid tin salt, metal salt of the various above mentioned carboxylic acid can be used conveniently

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carboxylic acid cobalt, a carboxylic acid zirconium, and carboxylic acid cerium, From a high point, the carboxylic acid titanium, and a carboxylic acid zirconium are still more preferred, and carboxylic acid Carboxylic acid potassium, carboxylic acid barium, carboxylic acid manganese, carboxylic acid nickel, acid vanadium, carboxylic acid iron, carboxylic acid titanium, carboxylic acid potassium, Carboxylic activity of a catalyst is preferred and Carboxylic acid bismuth, carboxylic acid calcium, Carboxylic Carboxylic acid calcium, carboxylic acid vanadium, carboxylic acid iron, carboxylic acid titanium, Carboxylic acid bismuth, carboxylic acid calcium, carboxylic acid vanadium, carboxylic acid iron, bismuth, carboxylic acid iron, and carboxylic acid titanium are especially the most preferred. in carboxylic acid metal salt other than said carboxylic acid tin salt, carboxylic acid bismuth, acid barium, carboxylic acid manganese, and a carboxylic acid zirconium are more preferred,

carboxylic acid potassium, carboxylic acid barium, carboxylic acid manganese, carboxylic acid nickel, material and weatherability which are obtained are high, and carboxylic acid bismuth, carboxylic acid calcium, carboxylic acid titanium, carboxylic acid potassium, carboxylic acid barium, and a carboxylic Carboxylic acid bismuth, carboxylic acid calcium, carboxylic acid vanadium, Carboxylic acid titanium coloring of a hardenability constituent obtained, and a point that the heat resistance of a hardened carboxylic acid cobalt, and a carboxylic acid zirconium, It is more desirable from a point with little acid zirconium are still more preferred.

It is more preferred that it is metal salt of a point of the ease (workability, viscosity) of dealing with it of carboxylic acid metal salt to monocarboxylic acid. 0238]

Bi(OCOR) 3 (23)

As said monocarboxylic acid metal salt, it is general formula (23) – (35). :

Ca(OCOR) 2 (24)

V(OCOR) 3 (25)

Fe(OCOR) 2 (26)

Fe(OCOR) 3 (27)

FI(0COR) 4 (28) K(OCOR) (29)

Mn(OCOR) 2 (31) Ba(OCOR) 2 (30)

nickel(OCOR)₂ (32)

Co(OCOR) 2 (33)

Zr (0) (OCOR) 2 (34)

Ce(OCOR) 3 (35)

(The inside R of a formula is substitution or an unsubstituted hydrocarbon group, and may include a carbon carbon double bond.) Two RCOO-bases may be the same and may differ. Carboxylic acid metal salt expressed is preferred.

acid radical of various carboxylic acid tin salt illustrated as the aforementioned (G) ingredient can be As a carboxylic acid group of carboxylic acid metal salt other than said carboxylic acid tin salt, an mentioned.

ethylhexanoic acid vanadium (trivalent), 2-ethylhexanoic acid calcium (divalent), 2-ethylhexanoic acid From a viewpoint of the availability of a raw material, and compatibility, as an example of desirable ethylhexanoic acid zirconium (tetravalence), 2-ethylhexanoic acid cerium (trivalent), neo decanoic acid bismuth (trivalent), Neo decanoic acid iron (divalent), neo decanoic acid iron (trivalent), neo (divalent), 2-ethylhexanoic acid iron (trivalent), 2-ethylhexanoic acid titanium (tetravalence), 2potassium (univalent), 2-ethylhexanoio acid barium (divalent), 2-ethylhexanoic acid manganese carboxylic acid metal salt, 2-ethylhexanoic acid bismuth (trivalent), 2-ethylhexanoic acid iron (divalent), 2-ethylhexanoic acid nickel (divalent), 2-ethylhexanoic acid cobalt (divalent), 2-

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JP,4101632,B [DETAILED DESCRIPTION]

barium (divalent), Manganese oleate (divalent), oleic acid nickel (divalent), oleic acid cobalt (divalent), (tetravalence), Naphthenic acid vanadium (trivalent), calcium naphthenate (divalent), naphthenic acid (trivalent), oleic acid iron (divalent), oleic acid iron (trivalent), oleic acid titanium (tetravalence), oleic (trivalent), Naphthenic acid iron (divalent), naphthenic acid iron (trivalent), naphthenic acid titanium calcium (divalent), Neo decanoic acid potassium (univalent), neo decanoic acid barium (divalent), a acid vanadium (trivalent), Oleic acid calcium (divalent), oleic acid potassium (univalent), oleic acid decanoic acid titanium (tetravalence), Neo decanoic acid vanadium (trivalent), neo decanoic acid neo decanoic acid zirconium (tetravalence), Neo decanoic acid cerium (trivalent), bismuth oleate An oleic acid zirconium (tetravalence), oleic acid cerium (trivalent), naphthenic acid bismuth naphthenic acid nickel (divalent), cobalt naphthenate (divalent), a naphthenic acid zirconium potassium (univalent), Naphthenic acid barium (divalent), manganese naphthenate (divalent), (tetravalence), naphthenic acid cerium (trivalent), etc. are mentioned. 2-ethylhexanoic acid bismuth (trivalent) from a viewpoint of catalytic activity, 2-ethylhexanoic acid iron (divalent), 2-ethylhexanoic acid iron (trivalent), 2-ethylhexanoic acid itanium (tetravalence). Neo acid iron (trivalent), oleic acid titanium (tetravalence), naphthenic acid bismuth (trivalent), Naphthenic decanoic acid bismuth (trivalent), neo decanoic acid iron (divalent), neo decanoic acid iron (trivalent), Neo decanoic acid titanium (tetravalence), bismuth oleate (trivalent), oleic acid iron (divalent), Oleic acid iron (divalent), naphthenic acid iron (trivalent), and naphthenic acid titanium (tetravalence) are more preferred, and 2-ethylhexanoic acid iron (trivalent), neo decanoic acid iron (trivalent), and especially naphthenic acid iron (trivalent) are preferred.

(tetravalence), 2-ethylhexanoic acid calcium (divalent), 2-ethylhexanoic acid potassium (univalent), 2acid zirconium (tetravalence), Naphthenic acid bismuth (trivalent), naphthenic acid ttanium (tetravalence), calcium naphthenate (divalent), naphthenic acid potassium (univalent), naphthenic acid decanoic acid zirconium (tetravalence), Bismuth oleate (trivalent), oleic acid titanium (tetravalence), 2-ethylhexanoic acid bismuth (trivalent) from a viewpoint of coloring. 2-ethylhexanoic acid titanium ethylhexanoic acid barium (divalent), 2-ethylhexanoic acid zirconium (tetravalence), Neo decanoic oleic acid calcium (divalent), Oleic acid potassium (univalent), oleic acid barium (divalent), an oleic (divalent), neo decanoic acid potassium (univalent), Neo decanoic acid barium (divalent), a neo acid bismuth (trivalent), neo decanoic acid titanium (tetravalence), Neo decanoic acid calcium barium (divalent), and a naphthenic aoid zirconium (tetravalence) are more preferred [0243]

Foluenesulfonic acid, styrene sulfonic acid, etc. are raised as organic sulfonic acid.

Alkyl acid phosphate is -0-P (=0) OH. It is phosphoric ester containing a portion and alkyl acid phosphate as shown below is contained. An organic acid nature phosphoric ester compound is preferred in respect of compatibility and curing catalyst activity. An organic acid nature phosphoric ester compound is expressed with $_{(R^{20},0)}\, _h^-P(=0)$ (-OH) $_{3^{-h}}$ (in the inside h of a formula, 1 or 2, and R^{20} show an organic residue).

Below, it illustrates concretely.

 $(CH_3O)_2 - P \ (=O) \ (-OH) \ and \ (CH_3O) - P (=O) \ (-OH)_2$, $(C_2H_5O)_2 - P \ (=O) \ (-OH) \ and \ (C_2H_5O) - P (=O) \ (-OH)_3$, $(C_3H_7O)_2 - P \ (=O) \ (-OH) \ and \ (C_4H_9O) - P \ (=O) \ (-OH)_3$ and $(C_4H_9O)_2 - P \ (=O) \ (-OH)_3$ and $(C_4H_9O)_3 - P \ (=O) \ (-OH)_4$ and $(C_4H_9O)_4 - P \ (=O) \ (-OH)_5$ and $(C_4H_9O)_4 - P \ (-OH)_5$ and $(C_4H_9O)_4$ $(=0) \ (-OH)_{2}, \ (G_8H_{17}O)_{2} -P \ (=0) \ (-OH) \ and \ (G_8H_{17}O) -P \\ (=0) \ (-OH)_{2}, \ (G_{10}H_{21}O)_{2} -P \ (=0) \ (-OH) \ and \ (-OH)_{2} + ($ $(\mathsf{G_{10}H_{21}O}) - \mathsf{P}(=0) \; (-\mathsf{OH}) \; {}_2 \; (\mathsf{G_{13}H_{27}O}) \; {}_2 - \mathsf{P} \; (=0) \; (-\mathsf{OH}) \; \mathsf{and} \; (\mathsf{G_{13}H_{27}O}) - \mathsf{P}(=0) \; (-\mathsf{OH}) \; {}_2 \; (\mathsf{G_{16}H_{33}O}) \; {}_2 - \mathsf{P} \; (-\mathsf{OH}) \; {}_3 \; (-\mathsf{OH}) \; {}_4 \; (-\mathsf{OH}) \; {}_4 \; (-\mathsf{OH}) \; {}_4 \; (-\mathsf{OH}) \; {}_5 \; (-\mathsf{OH}) \;$ (=O) (-OH) and (C $_{16}$ H $_{33}$ O)-P(=O) (-OH) $_2$ and $_{(HO-O_6H_{12}O)2}$ -P (=O) (-OH) and (HO-C $_6$ H $_{12}$ O)-P(=O) (-OH) $_2$, (HO-C $_8$ H $_1$ 6O)-P (=O) (-OH) and (HO-C $_8$ H $_1$ 6O)-P(=O) (-OH) $_2$, {(CH $_2$ OH) (CHOH) O] $_2$ -P

(=0) (-0H), ((CH20H) (CH0H) 0]-P(=0) (-0H) $_2$, Although ((CH20H) (CH0H) $_2$ H40] $_2$ -P (=0) (-0H) and ((GH $_2$ OH) (GHOH) ${
m G}_2{
m H}_4$ O}–P(=O) (–OH) $_2$ etc. are raised, it is not limited to the above–mentioned Illustration substance.

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By carboxylic acid, carboxylic acid metal salt other than carboxylic acid tin salt, organic sulfonic acid, and alkył acid phosphate, activity is low, and when moderate hardenability is not acquired, an amine compound can be added as a co-catalyst.

As various amine compounds, the indicated various above-mentioned amine compounds can be used as a co-catalyst of carboxylic acid tin salt (G).

loadings of an amine compound exceed 20 weight sections, pot life may become short too much and weight sections, and a hardening reaction becomes fully difficult to advance. On the other hand, if preferred, A cure rate may become it slow that loadings of an amine compound are less than 0.01 As for loadings of said amine compound, about 0.01-20 weight sections are preferred to organic polymer 100 weight section of an ingredient (A1), and also its 0.1 to 5 weight section is more is not preferred from a point of workability.

As a metal system compound of non-tin, besides carboxylic acid metal salt other than said carboxylic compound, an organic boron compound, etc. are preferred from a point of activity, it is not limited to acid tin salt, An organic metallic compound containing 3B fellows and 4A group metal is raised, and although a titanate organic compound, an organoaluminium compound, an organic zirconium

cetra (2-ethylhexyl titanate), Chelate compound, such as titanium chelate, such as titanium alkoxides, such as triethanolamine titanate, titanium tetra acetylacetonato, titanium ethylacetoacetate, octylene As said titanate organic compound, tetraisopropyl titanate, Tetrabuthyl titanate, tetramethyl titanate, glycolate, and titanium lactate, etc. are raised.

butoxy aluminum diisopropylate and aluminum sec-butyrate. Aluminum chelate, such as aluminum tris As said organoaluminium compound, aluminum isopropylate, Aluminum alkoxides, such as mono secacetylacetonato, aluminumtrisethylacetoacetate, and diisopropoxy aluminum ethylacetoacetate, is

As said zirconium compound, zirconium tetra isopropanal POKISAIDO, Zirconium alkoxides, such as a zirconium tetra-n PUROPI rate and zirconium normal butyrate. Zirconium chelate, such as zirconium tetra acetylacetonato, zirconium monoacetyl acetonate, zirconium bisacetylacetonate, zirconium acetylacetonato bis-ethylacetoacetate, and zirconium acetate, is raised.

concomitant use with said amine compound or an alkyl-acid-phosphate compound since it is possible organoaluminium compound, an organic zirconium compound, an organio boron compound, etc., It is to improve activity, and more desirable in a viewpoint of adjustment of working life in hardenability desirable in a viewpoint which can reduce the amount of catalyst used especially according to Although **** concomitant use is also possible so, these titanate organic compounds, an and ordinary temperature in an elevated temperature.

range, it is not desirable. On the other hand, if loadings exceed this range, working life becomes short too much, and workability may worsen, and it is not desirable from a point of storage stability. (E) As amount of ingredient used, about 0.01-20 weight sections are preferred to ingredient (A1) 100 weight section, and also about 0.5-10 weight sections are preferred. Since a oure rate may become slow and a hardening reaction will become fully difficult to advance if loadings are less than this

The (E) ingredient can be used combining two or more sorts besides using it alone.

In this invention, a minute hollow body can be used as a (F) ingredient. While improving the workability (******* thixotropy) of a constituent notably as indicated to JP,H11-35923,A or JP,H11-310772,A if this minute hollow body is used, it is known that a weight saving of a constituent and low-costizing are possible. However, it is known that the stability of a hardened material of a hardenability

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constituent and endurance which are obtained will fall according to an addition of this minute hollow

The hardenability constituent which added a minute hollow body of the (F) ingredient by using an organic polymer which is an ingredient (A1) of this invention as a polymer component can maintain righly the stability of a hardened material and endurance which are obtained, improving workability (*******) notably

a very small hollow body (henceforth a balloon) which is the (F) ingredient of this invention is a hollow body by which a diameter was preferably constituted from material of minerals of 500 micrometers or functional filler" (CMC). (F) An ingredient in particular is not limited but it is [ingredient] usable in less or quality of organicity 1 mm or less as indicated, for example on "state-of-the-art art of a various kinds of publicly known balloons.

strength of a hardened material may fall if average particle density is less than this range, and average particle density exceeds this range on the other hand, a workability improvement effect may preferred that it is 0.03-0.7 g/cm 3 , and it is preferred that it is especially 0.1-0.5 g/cm 3 . If tensile As for average particle density of a balloon, it is preferred that it is 0.01– $1.0 \mathrm{g/cm}^3$, it is more

An inorganic system balloon is more preferred than a point of stability and endurance to an organic system balloon.

ZIRCONIUM SPHEES made from ZIRCOA, KUREKASU fair made from Kureha Chemicals and product car boss fair made from GENERAL TECHNOLOGIES are marketed as a carbon balloon. etc. on a non–silicic acid system balloon. As an example of these inorganic system balloons, as a milt BALLOON made from EMERSON&CUMING, CELAMIC GLASSMODULES made from PITTSBURGE CORNING, As GLASS BUBBLES made from 3M, FUJIBA lune made from Fuji SHIRISHIA Chemicals, FILLTE U.S.A, As an alumina balloon, as BW by Showa Denko K.K., and a zirconia balloon HOLLOW balloon, a win light by LiICHI Chemicals, As a SANKI light by Sanki Engineering Co., Ltd., and glass Chemicals, and fly ash balloons, CEROSPHERES made from PFAMARKETING, FILLITE made from balloon, fly ash balloons, etc. can illustrate an alumina balloon, a zirconia balloon, a carbon balloon, As said inorganic system balloon, can illustrate a silicio aoid system balloon and a non-silicic acid system balloon, and on a silicic acid system balloon. A milt balloon, perlite, glass balloons, a silica and a silica balloon, as Q-CEL by Asahi Glass Co., Ltd., SAIRISHIA made from Fuji SHIRISHIA balloons, KARUN by Nippon Sheet Glass Co., Ltd., The Sumitomo 3M oell star Z-28, MICRO

here is made to foam, after blending a thing containing a foaming agent, and is good also as a balloon can illustrate a saran balloon, a polystyrene balloon, a polymethacrylate balloon, a polyvinyl alcohol balloon, and a styrene acrylic balloon at a thermoplastic balloon. A balloon of thermoplastics which system balloon, On a thermosetting balloon, a phenol balloon, an epoxy balloon, and a urea balloon A balloon of thermosetting resin and a balloon of thermoplastics can be illustrated as said organic constructed the bridge can also be used. A balloon after foaming may be sufficient, and a balloon

EMERSON&CUMING, As a urea balloon, ECCOSPHERES VF-O made from EMERSON&CUMING, As a saran balloon, SARAN MICROSPHERES made from DOW CHEMICAL, Expancel made from Japanese Filament, the Matsumoto Yushi-Seiyaku Matsumoto microsphere, As a polystyrene balloon, DYLITE EXPANDABLE POLYSTYRENE made from ARCO POLYMERS, SX863 by Japan Synthetio Rubber As an example of these organic system balloons, as a phenol balloon, Union Carbide UCAR and PHENOLIC MICROBALLOONS, As an epoxy balloon, ECGOSPHERES made from

The above-mentioned balloon may be used alone, and two or more kinds may be mixed and it may be used. The surface of these balloons Fatty acid, fatty acid ester, rosin, What was processed in order

Co., Ltd. (P) are marketed by EXPANDABLE POLYSTYRENE BEADS made from BASF WYANDOTE,

and constructed type styrene acrylic acid balloon of a bridge

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stiffening a compound, these balloons are used in order to carry out a weight saving and to cut down agent, titanium coupling agent, aluminum cup ring agent, a polypropylene glycol, etc. can be used. to improve dispersibility and the workability of a compound by rosin acid lignin, a silane coupling Without spoiling pliability, and elongation and intensity among physical properties at the time of

As for the amount of balloon used, about 0.1-50 weight sections are preferred to ingredient (A1) 100 improvement effect may not be enough if loadings are less than this range, and loadings exceed this range, tensile strength of a hardened material may fall or stability and endurance may worsen. weight section, and also its about 0.5-30 weight sections are preferred. When a workability

At this invention, it is a general formula as a (G) ingredient (7).

- SiR5 (OR6) 3-c (7)

oasis expressed can be used. General formula (6) which is the (A4) ingredient of this invention about ndependently, respectively, and c shows 0, 1, or 2.) -- an aminosilane coupling agent which has a (c R^5 is an organic group of monovalence of the carbon numbers 1–20 independently among a formula, respectively, 3-c R^6 is an organic group of monovalence of the carbon numbers 2-20this (G) ingredient :

- Si(OR4) 3 (6)

silicon atom is 2 to 20, toxic high methanol is not contained in alcohol generated in connection with a urns into a hardenability constituent with little change of a cure rate. A reactive silicon group of this have a methoxy group as an alkoxy group combined with a silicon atom, a reactant high methoxy silyl group does not generate to a reactive silicon group of the (A4) ingredient. Therefore, a hardenability which has a basis expressed with (R^{4in formula} is the same as the above), it becomes a hardenability While having stability, endurance, and creep resistance outstanding by adding to an organic polymer constituent in which an outstanding adhesive property is shown. To a reactive silicon group of this (G) ingredient and the (A4) ingredient, Since a carbon number of an alkoxy group combined with a (G) ingredient. Even if an ester exchange reaction between reactive silicon groups of the (G) ingredient and the (A4) ingredient advances after mixing with the (A4) ingredient since it does not constituent containing the (G) ingredient and the (A4) ingredient is before and after storage, and nydrolysis reaction of a reactive silicon group when a hardenability constituent carries out condensation hardening, but becomes it with a constituent with high safety.

constituent, when it is considered as 1 liquid type, since said hardenability constituent which consists (G) Although it is usable as a many liquid [such as 1 liquid type and a two-component type,] type of an ingredient and a (A4) ingredient has a large effect which makes small especially change of a cure rate in storage order, it is preferred.

(G) An ingredient is a compound which has a reactive silicon group expressed with a general formula (7), and an amino group. As an example of a reactive silicon group expressed with a general formula (7), a triethoxy silyl group, a methyldi ethoxy silyl group, a dimethylethoxy silyl group, an ethyldiethoxy alkoxy group combined with a silicon atom of a reactive silicon group has preferred toxic ethoxy silyl pieces are preferred, and its three pieces are more preferred. The toxicity of alcohol generated in connection with a hydrolysis reaction and a viewpoint of a cure rate to a triethoxy sllyl group is the number of an alkoxy group combined with one silicon atom of a reactive silicon group, two or more silyi group, a triisopropoxy silyi group, a methyldi isopropoxy silyi group, etc. can be mentioned. An reaction, and its ethoxy silyl group is more preferred. From a viewpoint of a cure rate, as for the group from a viewpoint or isopropoxy silyl of alcohol generated in connection with a hydrolysis most preferred.

triisopropoxy silane, gamma-aminopropyl methyldiethoxysilane, gamma-(2-aminoethyl) aminopropyl triethoxysilane, gamma-(2-aminoethyl) aminopropyl triisopropoxy silane, gamma-(2-aminoethyl) aminopropyl methyldiethoxysilane, gamma-ureido propyl triethoxysilane, gamma-ureido propyl As an example of an ingredient, (G) gamma-aminopropyl triethoxysilane, gamma-aminopropyl

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derivative and a condensation reaction thing of the above–mentioned silane compound can also use gamma-[2-(2-aminoethyl) aminoethyl] aminopropyl triethoxysilane, can be mentioned. A denatured triethoxysilane, N-vinylbenzyl gamma-aminopropyl triethoxysilane, N.N'-bis/gamma-triethoxy silyl propyl)ethylenediamine, Amino group content Silang, such as bis(triethoxy silyl propyl)amine and triisopropoxy silane, gamma-ureido propylmethyl diethoxysilane, N~phenyl-gamma-aminopropyl triethoxysilane, N–benzyl-gamma-aminopropyl triethoxysilane, N–n–butyl-gamma-aminopropyl the above-mentioned silane compound as a (G) ingredient.

The (G) ingredient used for this invention is used in 0.1-10 copies to 100 copies of organic polymers of the (A4) ingredient. It is preferred to use it in 1-5 copies especially. The above-mentioned (G) ingredient may be used only by one kind, and may carry out two or more kind mixing use.

silicon group of the (A4) ingredient is late is small and I the drying effect is high, it is desirable. Since compound which has alkoxy silyl groups and does not contain an amino group as a dehydrator storing such as the drying effect, hardenability, availability, and the tension physical properties of a hardened by care of health at low temperature comparatively since an ester exchange reaction with a reactive various kinds of compounds can be used. Since [that a change in physical properties after a silicon consists of an ingredient and a (G) ingredient. Especially as said dehydrator, it is not restricted but a silicon compound which has the Tori alkoxy silyl groups and does not contain an amino group has the higher drying effect, it is preferred, and especially a silicon compound that has a trimethoxysilyl group and does not contain an amino group is preferred. Specifically, alkyltrialkoxysilane, such as vinyltrimetoxysilane, methyl trimetoxysilane, and phenyltrimethoxysilane, is preferred from points, (A4) A dehydrator may be added when using as a 1 liquid type constituent a constituent which material

At this invention, it is a general formula as a (H) ingredient (8). :

- SiR7 (OCH3) (OR8) 3-d-8 (8)

numbers 2-20 independently, respectively, d shows 0, 1, or 2 and e shows 1, 2, or 3.) However, 3-de>=0 shall be satisfied. An aminosilane coupling agent which has a basis expressed oan be used. formula, respectively, R⁸ of a 3-d-e individual is an organic group of monovalence of the carbon (d R⁷ is an organic group of monovalence of the carbon numbers 1-20 independently among a General formula (6) which is the (A4) ingredient of this invention about this (H) ingredient :

and a reactant high methoxy silyl group generates to a reactive silicon group of the (A4) ingredient. A basis expressed with (R^{4tin} formula is the same as the above), (H) An ester exchange reaction between hardenability constituent obtained as a result turns into a hardenability constituent of fast curability If it is recuperated beforehand, a hardenability constituent added to an organic polymer which has a a methoxy silyl group of an ingredient and a reactive silicon group of the (A4) ingredient advances, while having outstanding adhesive property, stability, endurance, and creep resistance.

ingredient and a (A4) ingredient, Since it changes with existence of a transesterification catalyst and (A4) ingredient, etc., are not generally decided, but as a transesterification catalyst, When it includes temperature service, the 10-30 ** thing for which it is recuperated comparatively one week or more its addition, ester exchange reaction activity of a reactive silicon group of the (H) ingredient and the an organic tin catalyst or 0.5 copy - about three copies of Ti system catalysts in a system, in a low is preferred, and it is preferred that more than a day recuperates itself in not less than 30 ** high (H) Desirable care-of-health conditions of said hardenability constituent which consists of an temperature service.

constituent, when it is considered as 1 liquid type, since said hardenability constituent which consists (H) Although it is usable as a many liquid [, such as 1 liquid type and a two-component type,] type of an ingredient and a (A4) ingredient has a remarkable change of a cure rate especially by care of

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more pieces are preferred, and its three pieces are more preferred. Therefore, a trimethoxysilyl group (H) An ingredient is a compound which has a reactive silicon group expressed with a general formula (8), and an amino group. As an example of a reactive silicon group expressed with a general formula (8), A trimethoxysilyl group, a methyl dimethoxy silyl group, an ethoxy for the number of an alkoxy group combined with one silicon atom of a reactive silicon group, two or methoxy silyl group, etc. can be mentioned. From a viewpoint of ester exchange reaction speed, as dimethoxy silyl group, a dimethyl methoxy silyl group, a diethyl methoxy silyl group, a diethoxy is the most preferred.

As an example of an ingredient, (H) gamma-aminopropyl trimethoxysilane, gamma-aminopropyl methyl aminoethyl) aminoethyl] aminopropyl trimethoxysilane, can be mentioned. A denatured derivative and gamma-(2-aminoethyl) aminopropyl ethoxy dimethoxysilane, gamma-ureido propyltrimethoxysilane, benzyl-gamma-aminopropyl trimethoxysilane, N-n-butyl-gamma-aminopropyl trimethoxysilane, Ngamma–trimethoxysilypropyl)ethylenediamine, bis(trimethoxysilylpropyl)amine, and gamma–[2–(2– gamma-ureido propylmethyl dimethoxysilane, N-phenyl-gamma-aminopropyl trimethoxysilane, Na condensation reaction thing of the above-mentioned silane compound oan also use the aboveaminopropyl methyl dimethoxysilane, gamma-(2-aminoethyl) aminopropyl ethyl dimethoxysilane, vinylbenzyl gamma-aminopropyl trimethoxysilane, Amino group content Silang, such as N,N'-bis dimethoxysilane, gamma-(2-aminoethyl) aminopropyl trimethoxysilane, gamma-(2-aminoethyl) dimethoxysilane, gamma-aminopropyl ethyl dimethoxysilane, gamma-aminopropyl ethoxy mentioned silane compound as a (H) ingredient. The (H) ingredient used for this invention is used in 0.1-10 copies to 100 copies of organic polymers of the (A4) ingredient. It is preferred to use it in 1-5 copies especially. The above-mentioned (H) ingredient may be used only by one kind, and may carry out two or more kind mixing use. In this invention, an epoxy resin can be used as a (I) ingredient. This epoxy resin has a function which raises stability, endurance, and creep resistance further while improving impact strength and tough nature of an organic polymer which are the (A4) ingredients of this invention.

epoxy resin, A glycidyl ether type epoxy resin of a bisphenol A propylene oxide addition, p-oxybenzoic epoxy resins or novolak type epoxy resin is raised. Ranges of a using rate of these epoxy resin (f) and of a /epoxy resin becomes less than 1/100 and a rate of (A4)/epoxy resin surpasses 100/1, intensity of an organic polymer hardened material will become insufficient. Since a desirable using rate changes As an epoxy resin used as a (I) ingredient of this invention, an epichlorohydrin bisphenol A type epoxy diaminodiphenylmethane system epoxy resin, a urethane modified epoxy resin, Various cycloaliphatictough nature, stability, endurance, and creep resistance becomes will be hard to be acquired if a rate epoxy-resin and N,N-diglycidyl aniline, N,N-diglycidyl o-toluidine, Although an epoxidation thing of an improving intensity of a hardened material of the (A4) ingredient, it is good to carry out 5–50 weight– glycidyl ether of tetrabromobisphenol A, Novolak type epoxy resin, a hydrogenation bisphenol A type weight ratio. (A4) If the improvement effect of impact strength of an epoxy resin hardened material, section use of the epoxy resin five to 100 weight section still more preferably especially one to 200 unsaturation polymer, etc. are illustrated at the time, such as glycidyl ether of polyhydric alcohol, a currently generally used is used, and it gets. What contains an epoxy group in [two] a molecule at resin, Fire retardancy type epoxy resins, such as epichlorohydrin bisphenol F type epoxy resin and least has high reactivity when hardening, and a hardened material is preferred from points -- it is reactive silicon group containing organic polymer (A4) are (A4)/epoxy resin =100 / 1 - 1/100 in a easy to build three-dimensional meshes of a net. As a still more desirable thing, bisphenol A type preferably one to 100 weight section to epoxy resin 100 weight section. On the other hand, when improving the shock resistance of an epoxy resin hardened material, flexibility, tough nature, peel polyalkylene glycol diglycidyl ether, and glycerin, Not a thing limited to these but an epoxy resin with uses of a hardenability constituent, etc., are not generally decided, but. For example, when strength, etc., it is good to carry out 5-100 weight-section use of the (A4) ingredient still more hydantoin type epoxy resin, and petroleum resin, at the time, such as triglycidyl isocyanurate, acid glycidyl ether ester typed epoxy resin, m-aminophenol series epoxy resin, A

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weight section to (A4) ingredient 100 weight section.

Naturally a hardening agent which makes a constituent of this invention harden an epoxy resin can be used together. As an epoxy resin hardener which can be used, there is no restriction in particular and Triethylenetetramine, tetraethylenepentamine, Diethylamino propylamine, N-aminoethyl piperidine, mclasses, such as isophoronediamine and amine end polyether, second class amines;2,4,6-tris(dimethyl Anhydrous carboxylic acid; alcohols; phenols; carboxylic acid, such as DODESHINIRU succinyl oxide, xylylene diamine, m∸phenylenediamine, diaminodiphenylmethane, diaminodiphenyl sulfone, The first pyromellitic dianhydride, and anhydrous KUROREN acid; although compounds, such as a diketone aminomethyl) phenol, The third class amines like tripropylamine, and the salts; polyamide resin; complex compound of aluminum or a zirconium, can be illustrated, it is not limited to these. A compounds. Phthalic anhydride, hexahydro phthalic anhydride, tetrahydro phthalic anhydride, imidazole-derivatives; dicyandiamides of these third class amines; Boron trifluoride complex an epoxy resin hardener currently generally used can be used. Specifically, for example hardening agent may also be independent or two or more sorts may be used together.

When using a hardening agent of an epoxy resin, the amount used is the range of 0.1 to 300 weight section to epoxy resin 100 weight section.

have moisture, it exists stably, and it is decomposed into primary amine and ketone by moisture, and epoxy resin. If ketimine is used, a 1 liquid type constituent can be obtained. As such ketimine, it can Ketimine can be used as a hardening agent of an epoxy resin. In the state where ketimine does not produced primary amine serves as a hardening agent of the room-temperature-curing nature of an obtain by a condensation reaction of an amine compound and a carbonyl compound.

hexamethylenediamine, p∹phenylene diamine, and p,p'∸biphenylene diamine, Multivalent amine, such as Polyalkylene polyamine, such as TORIECHIREN triamine and tetraethylenepentamine; Polyoxyalkylene diaminobutane, pentamethylene diamine, 2,4-diaminopentane, Diamine;1,2,3-triamino propane, such as propionaldehyde, n-butylaldehyde, isobutyraldehyde, diethylacetaldehyde, Aldehyde, such as a glyoxal and benzaldehyde; Cyclopentanone, Cyclio ketone, such as trimethyl cyclopentanone, cyclohexanone, series polyamine gamma-aminopropyl triethoxysilane, Aminosilanes [, such as N-(beta-aminoethyl)and trimethyl cyclohexanone; Acetone, Methyl ethyl ketone, methyl propyl ketone, methyl isopropyl ketone, Methyl isobutyl ketone, a diethyl ketone, dipropyl ketone, diisopropyl keton, Aliphatic series triamino benzene, tris(2-aminoethyl) amine, and tetra(aminomethyl) methane; Diethylenetriamine, acetylacetone, methyl acetoacetate, ethyl acetoacetate, dimethyl malonate, diethyl malonate, a ketone, such as dibutyl ketone and diisobutyl ketone; beta-dicarbonyl compound [, such as an compound for composition of ketimine, As an amine compound, for example, ethylenediamine, gamma-aminopropyl trimethoxysilane and N-(beta-aminoethyl)-gamma-aminopropyl methyl Although what is necessary is just to use a publicly known amine compound and a carbonyl propylenediamine, Trimethylene diamine, a tetramethylenediamine, 1,3-diaminobutane, 2,3dimethoxysilane,]; etc. are used, and it gets. As a carbonyl compound, acetaldehyde, malonic acid methylethyl, and dibenzoylmethane,]; etc. can be used.

glycidyl ester, such as styrene oxide; butyl glycidyl ether and allyl glycidyl ether, etc. Such ketimines weight-section use is carried out to epoxy resin 100 weight section, and the amount used changes When an imino group exists in ketimine, an imino group may be made to react to glycidyl ether; may be used independently, two or more kinds may be used together and used for them, 1–100 with kinds of an epoxy resin and ketimine.

silicic acid anhydride, hydrous silicic acids, and carbon black; Calcium carbonate, Bulking agents, such bentonite, ferric oxide, a zinc oxide, an active white, and hydrogenation castor oil; fibrous fillers, such Various bulking agents other than a minute hollow body of the (F) ingredient may be blended with a hardenability constituent of this invention. It is not limited especially as said bulking agent, but For example, fumes silica, sedimentation nature silica, Reinforcement nature bulking agents, such as a as magnesium carbonate, diatomite, calcination clay, clay, talc, titanium oxide, bentonite, organic

is asbestos, glass fiber, and a filament, are illustrated

nainly chosen from titanium oxide, calcium carbonate, magnesium carbonate, talc, ferric oxide, a zinc oxide, etc. is used in the range of 5 - 200 weight section to organic polymer (A) 100 weight section. carbonate, calcination clay, clay, an active white, etc. is used in the range of 1 - 100 weight section desirable result will be obtained if a bulking agent chosen from surface treatment detailed calcium Of course, these bulking agents may be used only by one kind, and may mix and use two or more To obtain a hardenability constituent with high intensity with these bulking agents. Mainly Fumes constituent which is size with low strength, A desirable result will be obtained if a bulking agent silica, sedimentation nature silica, a silicic acid anhydride, hydrous silicic acids, carbon black, A to organic polymer (A) 100 weight section. When elongation wants to obtain a hardenability

enlarged or a lot of bulking agents can be mixed if a plasticizer is used, using it together with a in a hardenability constituent of this invention, since elongation of a hardened material can be bulking agent, it is more effective.

Diethylene giyool dibenzoate, Giyool ester, such as pentaerythritol ester; Butyl oleate, Aliphatic series 4s this plasticizer, dioctyl phthalate, dibutyl phthalate, Phthalic ester, such as butylbenzyl phthalate; paraffins, can use it independently arbitrarily in a form of two or more kinds of mixtures. A desirable result will be obtained if the amount of plasticizers is used in 100 or less weight sections to organic polyester plasticizer, polypropylene glycols, such as polyester of epoxy plasticizer, dibasic acid and dihydrio alcohol, such as epoxidation linseed oil and epoxy stearic acid benzyl, and a derivative of polybutadiene, Butadiene Acrylonitrile, polychloroprene, polyisoprene, polybutene, and chlorinated trioctyl phosphate and phosphorio acid actyldiphenyl; Epoxidized soybean oil, Polyether, such as Dioctyl adipate, Aliphatic dibasic acid ester, such as succinic acid isodecyl and dibutyl sebacate; sster species, such as methyl acetyl ricinolate, Tricresyl phosphate, Phosphoric ester, such as those; Polly alpha-methylstyrene, Polystyrene, such as polystyrene; plasticizers, such as polymer (A) 100 weight section.

A polymeric plasticizer can be used. If a polymeric plasticizer is used, as compared with a case where polytetramethylene glycol, or these polyether polyol An ester group, polyether [, such as a derivative Sebacic acid, Dibasic acid and ethylene glycol, such as adipic acid, azelaic acid, and phthalic acid, A property (it is also called paintwork) at the time of applying an alkyd paint to this hardened material diethylene glycol, triethylene glycol, propylene glycol, A polyester plasticizer obtained from dihydric a low molecule plasticizer which is a plasticizer which does not contain a polymer component in a molecule is used, early physical properties are maintained over a long period of time, and drying can be improved. A vinyl-base polymer produced by polymerizing by various methods in a vinyl species of polyalkylene glycols, such as triethylene glycol dibenzoate and pentaerythritol ester, system monomer as an example of a polymeric plasticizer, Diethylene glycol dibenzoate, Ester methylstyrene,]., although polybutadiene, polybutene, polyisobutylene, butadiene acrylonitrile, polyethylene glycols, a polypropylene glycol, A hydroxyl group of polyether polyol, such as changed into an ether group etc.,]; --- polystyrene [, such as polystyrene and Polly alphaalcohol, such as dipropylene glycol; 500 or more molecular weights, Further 1000 or more polychloroprene, etc. are mentioned, It is not limited to these.

and weatherability, and a vinyl-base polymer are especially preferred. Also in a vinyl-base polymer, an obtained an acrylic-acid-alkyl-ester system monomer indicated to JP,2001–207157,A by continuation polyacrylic acid alkyl ester, are still more preferred. Its molecular weight distribution is narrow, since preferred. Polyether and a vinyl-base polymer are preferred. A heat-resistant point to compatibility acrylic polymer and/or an methacrylic system polymer are preferred, and acrylic polymers, such as nypoviscosity—izing is possible for a synthetic method of this polymer, it is preferred, and it is still polymerization method] It is preferred to use a polymer what is called by a SGO process which Among these polymeric plasticizers, a polymer of the (A) ingredient and a thing to dissolve are more preferred. [of an atom-transfer-radical-polymerization method] [of a living-radical-

mass polymerization with an elevated temperature and high voltage.

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preferably. It is 1000-3000 most preferably. If a molecular weight is too low, a plasticizer can flow out plasticizer is not limited, a narrow thing is preferred and less than 1.80 are preferred. 1.70 or less are time, and alkyd paintwork cannot be improved. If a molecular weight is too high, viscosity will become although number average molecular weights of a polymeric plasticizer are 500–15000 preferably, they temporally by heat or a rainfall, early physical properties cannot be maintained over a long period of are 800-10000 more preferably — further — desirable — 1000-8000 — it is 1000-5000 especially high and workability will worsen. Although molecular weight distribution in particular of a polymeric more preferred, in addition, 1.60 or less are preferred, 1.50 or less are still more preferred, 1.40 especially or less are preferred, and 1.30 or less are the most preferred.

A number average molecular weight of a polymeric plasticizer and molecular weight distribution (Mw/Mn) are measured by the GPC method (polystyrene conversion).

Although a polymeric plasticizer does not have a reactive silicon group, it may have a reactive silicon group. When it has a reactive silicon group, it acts as a reaction plasticizer and shift of a plasticizer group, the number average molecular weight needs to be lower than a polymer of the (A) ingredient. molecule and one or less piece and 0.8 more piece or less are preferred. When using a plasticizer which has a reactive silicon group, especially an oxyalkylene polymer which has a reactive silicon from a hardened material can be prevented. When it has a reactive silicon group, it averages per

and a polymerio plasticizer may be used together. These plasticizers can also be blended at the time A plasticizer may be used alone and may use two or more sorts together. A low molecule plasticizer of polymer manufacture.

section preferably five to 150 weight section to (A) ingredient 100 weight section. In less than five The amount of plasticizer used is 20 - 100 weight section still more preferably ten to 120 weight weight sections, if an effect as a plasticizer stops being revealed and 150 weight sections are exceeded, mechanical strength of a hardened material runs short.

hydrocarbon group of the carbon numbers 1-20 independently among a formula, respectively.) a is 0, 1, 2, or 3. A silicon compound shown may be added. Although limitation is not carried out, as said hardenability constituent of this invention. $R_a Si(OR)_{4-a}$ (R is substitution or an unsubstituted It is a general formula in order to improve the activity of a condensation catalyst more in a silicon compound Phenyltrimethoxysilane, Phenylmethyldimethoxysilane,

phenyldimethylmethoxysilane, Since the effect that what is an aryl group of the carbon numbers 6~20 more preferred. If loadings of a silicon compound are less than this range, an effect of accelerating a diphenyldimethoxysilane, diphenyl diethoxysilane, and triphenylmethoxysilane, is preferred. Especially diphenyldimethoxysilane and diphenyl diethoxysilane are low oost, and especially since they are easy sections are preferred to (A) ingredient 100 weight section, and its 0.1 - 10 weight section is still hardening reaction may become small. On the other hand, when loadings of a silicon compound to receive, they are preferred. As for loadings of this silicon compound, about 0.01-20 weight accelerates a hardening reaction of a constituent is large, R in general formulas, such as exceed this range, hardness and tensile strength of a hardened material may fall.

vinyldimethylmethoxysilane, gamma-aminopropyl trimethoxysilane, The alkoxysilane; silicone varnishes alkoxysilane, such as dimethyldimethoxysilane, trimethylmethoxysilane, and n-propyltrimethoxysilane; Dimethyldi iso propenoxysilane, Alkyl iso propenoxysilane, such as methyl TORIISO propenoxysilane A physical-properties regulator which adjusts the tractive characteristics of a hardened material which have functional groups, such as N-(beta-aminoethyl) aminopropyl methyl dimethoxysilane, generated if needed to a hardenability constituent of this invention may be added. Although not limited especially as a physical-properties regulator, for example Methyl trimetoxysilane, Alkyl methyldimethoxysilane, gamma-glycidoxypropyltrimetoxysilane, Vinyltrimetoxysilane, and gamma-glycidoxy propylmethyl JIISO propenoxysilane, gamma-glycidoxy propyl

constituent of this invention is raised, or hardness is lowered conversely and elongation after fracture colysiloxanes are mentioned. By using said physical-properties regulator, hardness when stiffening a can be taken out. The above-mentioned physical-properties regulator may be used independently, gamma-mercapto propyltrimethoxysilane, and gamma-mercaptpropylmethyl dimethoxysilane; and may be used together two or more sorts.

especially a trimethyl silanol is preferred. A compound indicated to JP,H5-117521,A can be raised as without worsening stickiness of the surface of a hardened material. A compound which generates a compound which generates a compound which has a univalent silanol group in intramolecular by nydrolysis. A compound which generates a silicon compound which is a derivative of alkyl alcohol, such as a hexanol, octanol, and decanol, and generates R,SiOH **, such as a trimethyl silanol, by ntramolecular by hydrolysis has the operation which reduces a modulus of a hardened material Especially a compound that generates a compound which has a univalent silanol group in

glycerin, pentaerythritol, or sorbitol, are three or more, and generates R3SiOH **, such as a trimethyl yydrolysis, Trimethylolpropane indicated to JP,H11-241029,A, A compound which generates a silicon compound which is a derivative of polyhydric alcohol whose numbers of hydroxyl groups, such as

silanol, by hydrolysis can be raised.

monosilanol content compound by a hydrolytic silicon content group and hydrolysis in which bridge A compound which generates a silicon compound which is a derivative of an oxypropylene polymer nydrolysis can also be raised. A polymer which has a silicon content group which can serve as a which is indicated to JP,H7-258534,A, and generates R₃SiOH(s), such as a trimethyl silanol, by construction furthermore indicated to JP,H6-279693,A is possible can also be used.

A physical-properties regulator is preferably used in the range of 0.5-10 weight section 0.1 to 20weight section to (A) ingredient 100 weight section.

These thixotropic grant agent (lappet inhibitor) may be used independently, and may be used together In a hardenability constituent of this invention, a lappet is prevented if needed, and in order to improve workability, a thixotropic grant agent (lappet inhibitor) may be added. Although not limited two or more sorts. A thixotropic grant agent is used in the range of 0.1-20 weight section to (A) derivative; calcium stearate, aluminum stearate, and barium stearate, is mentioned, for example. especially as a lappet inhibitor, metallic soap, such as polyamide wax; hydrogenation castor oil ingredient 100 weight section.

[0303]

epoxy compound is good to use it in the range of 0.5-50 weight section to (A) ingredient 100 weight A compound which contains an epoxy group in one molecule in a constituent of this invention can be used. If a compound which has an epoxy group is used, the stability of a hardened material can be acid ester, alicycle fellows epoxy compounds, and an epichlorohydrin derivative as a compound which epoxidation linseed oil, a di(2-ethylhexyl) 4,5-epoxy cyclohexane~1,2-JIKABOKISHI rate (E-PS), epoxy octyl stearate, epoxy butyl stearate, etc. are raised. Especially in these, E-PS is preferred. An mproved. Compounds shown in epoxidation unsaturation oil and fat, epoxidation unsaturation fatty has an epoxy group, those mixtures, etc. can be illustrated. Specifically, epoxidized soybean oil,

of a hardened material, and dust is carried out. Drying oil represented with tung oil, linseed oil, etc. by material, and an operation of preventing adhesion of stickiness of the surface, garbage on the surface polybutadiene produced by making carry out copolymerization, Liquefied polymers, such as a polymer compound; An acrylic polymer which denaturalized with drying oil, Epoxy system resin, silicon resin; example of an oxygen hardenability substance, Various alkyd resins produced by denaturalizing this An oxygen hardenability substance can be used for a constituent of this invention. To an oxygen Butadiene, chloroprene, isoprene, Diene series, such as 1,3-pentadiene, a polymerization or 1,2hardenability substance, an unsaturated compound which can react to oxygen in the air can be llustrated, it reacts to oxygen in the air, a cured film is formed near the surface of a hardened

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these, tung oil and a liquefied diene system polymer are preferred. Concomitant use of a catalyst and weight section. If an improvement of stain resistance becomes less enough when said amount used tractive characteristics of a hardened material, etc. to be spoiled will arise. An oxygen hardenability of 1,4-polybutadiene, C5 - C8 diene, NBR produced by making carry out copolymerization of these diene series may serve as a subject, Liquefied copolymers, those various denaturation things, etc. (catalysts and metal driers, metal salt, such as cobalt naphthenate, lead naphthenate, a naphthenic diene series and the monomers which have copolymeric, such as acrylonitrile and styrene, so that These may be used independently and may be used together two or more sorts. Especially among ingredient 100 weight section, and the amount of oxygen hardenability substance used is 0.5-10mallein-ized denaturation thing, a boiled oil denaturation thing, eto.), such as SBR, are mentloned. illustrated. It is at best still more preferred to use it in the range of 0.1-20 weight section to (A)substance is good to use it, using together with a photoresist substance as indicated to JP,H3~ will be less than 0.1 weight sections, and 20 weight sections are surpassed, a tendency for the a metal drier which promote an oxidation hardening reaction may heighten an effect. As these acid zirconium, octylic acid cobalt, and an octylic acid zirconium, an amine compound, etc. are

A photoresist substance can be used for a constituent of this invention. If a photoresist substance is GURIKORUJI (meta) acrylate and neopentyl GURIKORUJI (meta) dimethacrylate, or with a molecular substance produces physical-properties change of hardening etc. Many things, such as a constituent partly, It is mixtures, such as oligomer or it, and monomers, such as propylene (or butylene, ethylene) used, a coat of a photoresist substance is formed in the hardened material surface, and stickiness of ARONIKKUSU M-210 (2 Functional), ARONIKKUSU M-215, ARONIKKUSU M-220, ARONIKKUSU Mcontaining especially an acrylic functional group is preferred, and a compound which averages in one light, molecular structure causes a chemical change considerably for a short time, and a photoresist ARONIKKUSU M-369, ARONIKKUSU M-310, Although ARONIKKUSU M-315, ARONIKKUSU M-320, a hardened material and the weatherability of a hardened material can be improved. By operation of compound, acrylic or an methacrylic system unsaturation group 1 thru/or a monomer which it has containing an organic monomer, oligomer, resin, or them, are known by this kind of compound, and 233, ARONIKKUSU M-240, ARONIKKUSU M-245; (three organic functions) ARONIKKUSU M-305, molecule and contains the three or more functional groups is preferred. (Each ARONIKKUSU is a ARONIKKUSU M-325, (polyfunctional) ARONIKKUSU M-400, etc. can be illustrated, a compound compound, polycinnamic acid vinyl, or azide-ized resin can be used. As an unsaturation acrylic weight of 10,000 or less oligoésters is illustrated. Specifically, For example, special acrylate. commercial arbitrary things can be adopted as it. As a typical thing, an unsaturation acrylic product of Toagosei chemical industry incorporated company above.)

and amines, may heighten an effect. A photoresist substance is good to use it in the range of 0.5-10as a sensitization group, usually, a "photopolymer" (Showa 47(1972) — on March 17) besides [which illustration — these — a sensitizer can be used, being able to mix and adding [it can be independent, A polycinnamic acid vinyl derivative of many besides what is a photopolymer which uses a cinnamoyl weight section preferably 0.1 to 20 weight section to (A) ingradient 100 weight section, and in 0.1 or less weight section, since there is no effect which improves weatherability, and a hardened material cinnamic acid is illustrated. Azide-ized resin is known as a photopolymer which uses an azido group or] if needed. Addition of accelerators, such as sensitizers, such as ketone and a nitro compound, becomes hard too much and produces a cracking crack in 20 or more weight sections, it is not added a diazido compound as a sensitizing agent] a rubber sensitizing solution [and] printing group as a sensitization group as polycinnamic acid vinyl, and esterified polyvinyl alcohol with society publication part issue, and the 93rd page – 106th page – 117th page – have detailed

used, the weatherability of a hardened material oan be improved. Although a hindered phenol system, An antioxidant (antiaging agent) can be used for a constituent of this invention. If an antioxidant is antioxidant, especially a hindered phenol system is preferred. Similarly, Tinuvin 622LD, tinuvin 144; CHIMASSORB944LD, CHIMASSORB119floor line (all are the Ciba~Geigy Japan, Inc. make above); a mono- phenol system, a bisphenol system, and a polyphenol system can be illustrated as an

MARK LA-57, MARK LA-62, MARK LA-67, and MARK. LA-63 and MARK. LA-68. (All are ADEKAAGASU chemicals incorporated company make above). A hindered amine light stabiliser shown in SANORU LS-770, SANORU LS-7765, SANORU LS-292, SANORU LS-2626, SANORU LS-1114, and SANORU LS-744 (all are the Sankyo Co., Ltd. make above) can also be used. An example of an antioxidant is indicated also to JP,H4-283259,A or JP,H9-194731,A. It is at best still more preferred to use it in the range of 0.1 – 10 weight section to (A) ingredient 100 weight section, and the amount of antioxidant used is 0.2 – 5 weight section.

Light stabilizer can be used for a constituent of this invention. If light stabilizer is used, photooxidation degradation of a hardened material can be prevented. Although a benzotriazol system, a hindered amine system, a benzoate system compound, etc. can be illustrated as light stabilizer, especially a hindered amine system is preferred. It is at best still more preferred to use it in the range of 0.1 - 10 weight section to (A) ingredient 100 weight section, and the amount of light stabilizer used is 0.2 - 5 weight section. An example of light stabilizer is indicated also to JP,H9-194731,A.

When an unsaturation acrylic compound is used especially as a photoresist substance in a constituent of this invention, it is preferred to use a tertiary amine content hindered amine light stabiliser as a hindered amine light stabiliser as indicated to JP,H5-70531,A because of preservation stability improvement of a constituent. As a tertiary amine content hindered amine light stabiliser. ** tinuvin 622LD and tinuvin 144;. CHIMASSORB119floor line. (All are the Ciba-Geigy Japan, Inc. make above);MARKLA-57, LA-62, LA-67, LA-63 (all are ADEKAAGASU chemicals incorporated company make above); SANORU LS-765, LS-292, LS-2626, LS-1114, LS-744. (All are the Sankyo Go., Ltd. make above) etc. — light stabilizer can be illustrated.

An ultraviolet ray absorbent can be used for a constituent of this invention. If an ultraviolet ray absorbent can be used for a constituent of this invention. If an ultraviolet ray absorbent is used, the surface weatherability of a hardened material can be improved. Although a benzophenone series, a benzotriazol system, a metal chalet esystem compound, etc. can be illustrated as an ultraviolet ray absorbent, especially a benzotriazol system is preferred the same tail more preferred to use it in the range of 0.1 – 10 weight section to (A) ingredient 100 weight section, and the amount of ultraviolet ray absorbent used is 0.2 – 5 weight section. It is preferred to use together and use a phenol system, a hindered phenolic antioxidant, a hindered amine light stabiliser, and a benzotriazol system ultraviolet ray absorbent

An ingredient which limitation in particular does not have in the method of preparation of a hardenability constituent of this invention, for example, was described above is blended, it kneads under ordinary temperature or heating using a mixer, a roll, a kneader, etc., or an ingredient is dissolved using a little suitable solvents, a usual method of mixing is adopted, and it gets. A many liquid L such as 1 liquid type and a two-component type, I type compound can also be made and used by combining these ingredients suitably.

by the hardenability constituent of this invention is exposed into the atmosphere, by operation of moisture, it will form network structure in three dimensions, and will harden it promptly to a solid which has rubber-like elasticity.

It faces using a hardenability constituent of this invention, if needed Adhesive improving agents other than an aminosilane, a physical-properties regulator. It is possible to add suitably various additive agents, such as a preservation stability improving agent, an ultraviolet ray absorbent, a metal deactivator, anti-ozonant, light stabilizer, amine system radical chain inhibitor, the Lynn system peroxide decomposition agent, lubricant, paints, and a foaming agent.

A hardenability constituent of this invention can be used for sealant, such as a binder, a building, a marine vessel, and a super highway, adhesives, modeling material, a vibroisolating material, a sound deadener, a sound insulating material, a charge of foam, a paint, a gunning material, etc. Electrical insulation materials, such as electric electronic component materials, such as a solar cell rear-face sealing agent, pre-insulation an electric wire, material for cables, Elastic adhesives, powder coatings,

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large area, such as glass, porcelain, wood, metal, and a resin-molding thing, it is usable also as various proof of the glass laminate end face (cut section), autoparts, electrical machinery parts, several kinds material for masonry joints of sheathing materials, such as a medical equipment sealant, food packing electric electrons, a film, a gasket, It is available for various uses, such as a fluid-scalant agent used from excelling in stability, endurance, and creeping property. Adhesives for interior panels, adhesives finishing adhesives, adhesives for finishing of wall, adhesives for car panels, It is desirable, especially of machine part, etc. Since, or help of a primer is borrowed and it may stick to substrates of a **** material, and a sizing board, A coating material, a primer, a conductive material for electromagnetic precision-mechanical-equipment assemblies, a sealing material for direct grazing, a sealing material seal constituents and adhesion constituents of a type. A hardenability constituent of this invention for face panels, adhesives for tiling, adhesives for stone tensions, Ceiling finishing adhesives, floor for multiple glass, a sealing material for speed signal generator construction methods, or a sealing in various molding materials and wired sheet glass and a sealing agent for rust prevention / water wave cover, a thermally conductive material, A charge of a hot melt material, a potting agent for casting material, a medical-application rubber material, a medical-application binder, A sealing when it is considered as the electrical and electric equipment, an electron and adhesives for material for working joint of a building and uses.

[0314]

[Example] Although working example is hung up over below and this invention is explained to it in more detail, this invention is not limited only to these working example.

(Synthetic example 1)

Use polyoxypropylene triol of the molecular weight 3,000 [about] as an initiator, and propylene oxide is polymerized in a zinc hexa cyanocobaltate glyme complex compound catalyst, Number average molecular weight about 26,000 (polystyrene reduced molecular weight in which the column measured the solvent using THF using the TOSOH TSK-GEL H type using TOSOH HLC-8120GPC as a liquid-sending system) polypropylene oxide was obtained. Then, the methanol solution of NaOMe of the equivalent was added 1.2 times to the hydroxyl group of this hydroxyl group end polypropylene oxide, and methanol was distilled off, and also the allyl chloride was added, and the hydroxyl group of the end was changed into the allyl group. Decompression devolatilization removed the unreacted allyl chloride. To aliy end polypropylene oxide 100 weight section which is not refined [which was obtained], n-hexane 300 weight section. After it carried out mixed stirring of the water 300 weight section further at the hexane solution obtained by centrifugal separation removing water after carrying out mixed stirring of the water 300 weight section and centrifugal separation removed water again, decompression devolatilization removed hexane. By the above, the end obtained 3 organiofunctions polypropylene oxide of the number average molecular weight 26,000 [about] which is an allyl group.

150 ppm of platinum content 3wt% of platinum vinyl siloxane complexes isopropanol solutions are made into a catalyst to allyl end polypropylene oxide 100 obtained weight section, it was made to react to methyl dimethoxysilane 1.4 weight section at 90 ** for 5 hours, and the methyl dimethoxy silyl group end polyoxyalkylene series polymer (A-1) was obtained. Measurement by 1H-NMR (it measures in a CDCl₃ solvent using JEOL, JNM-LA400) averaged the methyl dimethoxy silyl group of

the end per molecule, and they were 2.3 pieces.

(The reference example 1, working example 2-4, and comparative examples 1-2)
Organic polymer (A-1) 100 weight section which has the reactive silicon group obtained in the synthetic example 1 according to the combination formula shown in Table 1. Surface treatment colloid calcium carbonate (product made from Shiraishi industry, Hakuenka GGR) 120 weight section, Titanium oxide (Ishihara Sangyo make, TIPAQUE R-820) 20 weight section, DIDP55 weight section, the amount part of thixotropic grant agent (made in [Kusumoto Chemicals], DISUPARON 6500) duplexs, and light stabilizer (the Sankyo make, SANORU LS7701 weight section and an ultraviolet ray absorbent (made in Tiba Specialty Chemicals.) Tinuvin 3271 weight section, antioxidant (product

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made from Ouchi Shinko Chemical industry, NOKURAKKU SP) 1 weight section, The amount part of

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the product made from neo decanoic acid tin (divalent) (trade name: U-50); Japan epoxy resin.) given the dibutyltin bisaoetylacetonate (trade name: U-220), Japanese east --- transformation --- make and (aminoethyl)-gamma-aminopropyl trimethoxysilane (the Nippon Unicar make.) A-1120) silicate (made dehydrator vinyltrimetoxysilane (Nippon Unicar make, A-171) duplexs, adhesion grant agent N-betain a col coat.) given in three weight sections and Table 1 Made in an ethyl silicate 28; col coat, ethyl silicate 40; Made in a col coat, a curing catalyst (the Japanese east -- transformation -- make and in the amount part of methylsilicate 51 duplexs, and Table 1 Neo decanoic acid (trade name; BASA substantially under drying conditions, it sealed in the dampproof container and 1 liquid mold-curing carried out number-of-copies addition, after kneading in the state where moisture does not exist tick 10); the description to Table 1 of the Wako Pure Chemical Industries make and lauryl amine nature constituent was obtained.

(Hauling physical properties of a hardened material)

It was 23 **x3 +50 **x recuperated in the class product of Table 1 on the 4th, and the sheet about 3 mm thick was created. [per day] It examined by having pierced this sheet to the No. 3 dumbbell type, and having pulled by a part for 200-mm/in hauling speed, and was extended at the time of intensity (MPa) and Eb:fracture at the time of M50:50% hauling modulus (MPa) and Tb:fracture, and (%) was measured. A result is shown in Table 1.

Recovery)

It was 23 **x3 +50 **x recuperated in the class product of Table 1 on the 4th, and the sheet about 3 mm is pulled to 40 mm between the marked lines (100% extension), it fixed at 60 ** for 24 hours. The recovery was measured from the rate which opened this wide at 23 ** and the marked line restored I hour afterward. It means that the one where the recovery is larger is excellent in stability. A result mm thick was created. [per day] This sheet was pierced to the No. 3 dumbbell type, and where 20 is shown in Table 1.

(Creep measurement using the piece of a dumbbell)

It was 23 **x3 +50 **x recuperated in the class product of Table 1 on the 4th, and the sheet about 3 mm thick was created. [per day] This sheet was pierced to the No. 3 dumbbell type, and the marked line of 20 mm of intervals was described. The end of this piece of a dumbbell was fixed in 60 ** oven, where a displacement difference is smaller is excellent in creep resistance. A result is shown in Table marked lines of 200 hours after immediately after imposing load was measured. It means that the one lower end of the hung piece of a dumbbell. The displacement difference of the distance between the mentioned tension physical-properties measurement of this hardened material was imposed on the and the piece of a dumbbell was hung. 0.4 time as much load as M50 value obtained by the above-

Table 1

2 1 4 8 8 9 88 88 88 88 88 88 88 88 88 88 88									
株式 大型 大型 大型 大型 大型 大型 大型 大	081	382	389	745	400	Z6Z	(%)	q∃	
株式 大型 大型 大型 大型 大型 大型 大型 大			2.35	2.61	72.27	2.00	(sqM)		
2 1 4 8 8 88 88 88 88 88 88 88 88 88 88 88		0.42	09.0	84.0	97 0	0.53	(BPA)	0 9 M	樹子宮宮
2 1 4 2 3 4 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	2.2	30.2	7 'l	1.2	9.1	1.61			
2 1 本 8 2 1 1 0 1 001 001 001 001 001 001 001 001 0	08	97	98	88	1/8	7 9	(%)		彰
1	97.0		9L '0	<i>9L</i> '0	97.0		くきてみいたて	Ć.	
1			1.2	1.2	2.1		01467 14 - V		
2 1 4 8 8 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	3.4		3.4	3.4	3.4		03-U<&X\\		
2 1 4 8 2 1 2 3 4 1 2 3 4 1 2 3 4 1 3 4 9 2 3 4 9 2 3 4 9 2 3 4 9 2 3 4 9 2 3 4 9 2 3 4 9 2 3 4 9 2 3 4 9 2 3 4 9 2 3 4 9 2 3 4 9 2 3 4 9 2 3 4 9 2 3 4 9 2 3 4 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 2 3 9 9 9 2 3 9 2 3		7				7	447470-220	卓機XX.	硬化触媒
2 1 4 2 3 4 1 2 3 4 1 2 3 4 1 3 4 1 3 4 1 3 4 1 3 4 1 3 4 1 3 4 1 3 4 1 3 4 1 3 4 1 3 4 2 3 3 4 3 3 3			7						
2 1 4 8 2 1 2 1 4 6 6 7 1<				7					
2 1 4 8 2 1 001 0					7		824-411<14I		1-40%
2 1 4 8 2 1 00 10	3	3			3	3	0211-A		
2 1 4 8 2 1 001 0	7	7	7	7	7	7			
2 1 4 8 2 1 001 0	Ţ.	ļ	ļ	1	1	ļ			
2 1 4 8 2 1 001 0	Į.	L	Ţ	L	I	l.	<u>+</u> 35, >357		
2 1 4 5 2 1 4 1 6 1<	ļ	1	ļ	1	L	l l	0 <i>LL</i> -S74-/4		
2 1 2 3 4 1 2 1 4 4 1 3 4 1 1 3 4 1<	7	7	7	7	7	7	0039# <pre>'\X\'T</pre>		
2 1 4 8 2 1 </td <td>99</td> <td>99</td> <td>99</td> <td>99</td> <td>99</td> <td>99</td> <td></td> <td>トトラー </td> <td><u>[п</u></td>	99	99	99	99	99	99		トトラー	<u>[п</u>
2 I A E 2 I 001 001 001 001 100 100 100 100 100 10	50	50	70	70	70	50	₽4√° −9R−820		
7 1 7 3 4 1 2	120	120	120	120	120	120	800華體白		
	001	100	100	100	100	100	- Α	会 類(A)	朴合重辦育
「松本の」 「松本」 「松本冬」 (信重単) 刈邸	7	Ţ	7	_	7	L			
内 ・	科技	≢ 77		例就実		例等卷		(陪量重) 丸 的	

silicate. As shown in the comparative example 2, when carboxylic acid tin salt (neo SUTAN U-50) etc. shown in the reference example 1, stability and creep resistance are notably improved by addition of are used, not using organic tin (U-220) as a curing catalyst, stability and creep resistance also with catalyst, especially the recovery of creep resistance is low bad silicate additive-free. However, as As shown in the comparative example 1 of Table 1, when organic tin (U-220) is used as a curing good silicate additive-free are shown, but. As shown in working example 2-4, stability and creep resistance further outstanding by silicate addition were shown. The ethyl silicate 40 and the http://www4.ipdl.inpit_go.jp/cgi=bin/tran_web_cgi_ejje?atw_u=http%3A%2F%2FwzFwww4.ipdl.i.. 2010/04/30

methylsilicate 51 which were used in working example 3-4 are a condensate of a tetraethoxysilane and a tetramethoxy silane, respectively, and showed the especially outstanding effect

(Synthetic example 2)

group end polypropylene oxide of the number average molecular weight 14,500 [about] produced by polymerizing propylene oxide in the zinc hexa oyanocobaltate glyme complex compound catalyst is To this allyl end polypropylene oxide, in the same procedure as the synthetic example 1, it was made Use polyoxypropylene glycol of the molecular weight 2,000 [about] as an initiator, and the hydroxyl used, Allyl end polypropylene oxide was obtained in the same procedure as the synthetic example 1. to react to trimethoxysilane and the polyoxyalkylene series polymer (A-2) which has an average of .5 trimethoxysilyl groups at the end was obtained.

(Synthetic example 3)

To the allyl end polypropylene oxide obtained in the synthetic example 2, in the same procedure as the synthetic example 1, it was made to react to triethoxysilane and the polyoxyalkylene series colymer (A-3) which has an average of 1.5 triethoxy silyl groups at the end was obtained

(Synthetic example 4)

To the ally end polypropylene oxide obtained in the synthetic example 2, in the same procedure as the synthetic example 1, it was made to react to methyl dimethoxysilane and the polyoxyalkylene series polymer (A-4) which has an average of 1.5 methyl dimethoxy silyl groups at the end was obtained.

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(The reference examples 5-11 and comparative examples 3-5)

the amount part of thixotropic grant agent (made in [Kusumoto Chemicals], DISUPARON 6500) duplexs, and light stabilizer (the Sankyo make.) SANORU LS7701 weight section and an ultraviolet ray absorbent (made in Tiba Specialty Chemicals.) Tinuvin 3271 weight section, antioxidant (product colloid calcium carbonate (product made from Shiraishi industry, Hakuenka CCR) 120 weight section, U-220), the Sankyo Organic Chemicals make, a dibutyltin JIRAURI rate (trade name: STANN BL)) of nade from Ouchi Shinko Chemical industry, NOKURAKKU SP) 1 weight section, The amount part of dehydrator vinyltrimetoxysilane (Nippon Unicar make, A-171) duplexs, adhesion grant agent N-betaaminoethyl}-gamma-aminopropyl trimethoxysilane (the Nippon Unicar make.) A-1120) Three weight name: U-50) and amine (the Wako Pure Chemical Industries make, lauryl amine) carried out number-Titanium oxide (Ishihara Sangyo make, TIPAQUE R-820) 20 weight section, DIDP12 weight section, synthetic examples 2-4 according to the combination formula shown in Table 2, Surface treatment curing catalyst (Japanese east transformation make and dibutyltin bisacetylacetonate (trade name: drying conditions, it sealed in the dampproof container and 1 liquid mold-curing nature constituent sections, number of copies given [silicate (made in a col coat, methylsilicate 51)] in Table 2, the of-copies addition, after kneading in the state where moisture does not exist substantially under Organic polymer (A-2-4) 100 weight section which has the reactive silicon group obtained in the make.) of the (C) ingredient The description to Table 2 of neo decanoic acid tin (divalent) (trade the (D) ingredient given in Table 2, or the curing catalyst (the Japanese east — transformation vas obtained.

examined by having pulled by the same method as the above-mentioned using the class product of Table 2, and was extended at the time of intensity (MPa) and Eb:fracture at the time of M50:50% hauling modulus (MPa) and Tb:fracture, and (%) was measured. A result is shown in Table 2. The recovery was measured by the same method as the above-mentioned using the class product of recovery was measured from the rate which opened this wide at 23 ** and the marked line restored 1 hour afterward. A result is shown in Table 2. Table 2. However, the stretched state was fixed at 23 ** 100% for 24 hours this time, and the

(Creep measurement using a shear sample)

The displacement difference with the 140-hour back immediately after creating the ** sample which

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0.4 mm, and the displacement difference carried out a thing of 0.4 mm or more. A result is shown in is not carried an area of 20 mm x 25 mm and 1 mm in thickness using the class product of Table 2, imposing load was measured. [per day] The displacement difference made θ x for the thing below imposing 0.1MPa load for what recuperated [23 **x3 +50 **x] itself on the 4th in 60 ** oven, and Table 2.

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Table 2

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370	340	697	516	<u> </u>	802	802	18t	961	193	(%)	9 🗎	
2, 70	78.2	2, 83	81.2	2.20	80.2	2.35	7.61	2, 20	2, 24	(s¶M)	dТ	
0.82	98.0	10.1	ħ6 ˙0	۱. او	76.0	76 '0	1 13	96 '0	76 0	(RPA)	OSM	種化物物体
×	×	×	0	0	0	0	0	0	0		(根公野)	
7 9	ΖÞ	<i>L</i> 9	96	96	†6	86	ε6	63	66	(%)	率元	彰
			91.0							くきていけたさ	マミヤ	
			3,4							09 -በ ረ <i>६</i> ሂ <i>ዩ</i> ኔ	(C) 就分	
3					3			0.3		Stann BL		
	7	2.0		7		7	0.2		2.0	022-U<₹ <u>₹</u>	会額(□)	類軸小動
				7						131-4(14)(4)	会熟(日)	イーセルぐ
3	3	3	3	3	3	3	3	3	8	0211~A	 	
7	7	7	7	7	7	7	7	7	7	171-A	(南水	
1	Į.		L	L	L	L	l.	l	L	d\$46 <u>£</u> 47	[廃土 初	
1	l l	Į.	I.	L	Ţ	1		l L	ļ	£₹£, >35 <u>7</u>	喀 观观	
l	L	L	l.	Ţ	l.	1	L L	l l	l l	0 <i>LL</i> -\$711-74	[隆宝]	
2	7	7	7	2	7	7	7	7	7	4, (YV. ti) #6500	性位与例	
71	15	15	12	15	15	15	15	15	15	9010	降壓	[0
20	50	50	50	50	50	50	50	50	50	44v4B-850		
150	120	150	120	150	150	150	120	150	150	AOO華醴白		莊
001	001	001								基心では、公は、カーA		
			100	100	100	100	100			基机於科扎州 E-A		
								100	100	基心で対化 3ーA	会類 (FA)	本合重 数字
g	7	3	ΙL	01	6	- 8	L	9	G	野なり		
	医棘 型					侧岩卷				素トイ卦③艮	(陪量重) 如服	¥

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When the reactive silicon group of an end uses the organic polymer (A–2–3) which is the Tori alkoxy examples 3–5, stability and creep resistance are improved notably. The reference example 10 which added silicate, and the reference example 11 using carboxylic acid tin salt (neo SUTAN U-50) as a silyl groups from comparison with the reference examples 5-9 of Table 2, and the comparative curing catalyst showed the further outstanding recovery.

(Synthetic example 5)

To the allyl end polypropylene oxide obtained in the synthetic example 1, in the same procedure as the synthetic example 1, it was made to react to methyl dimethoxysilane and the polyoxyalkylene series polymer (A-5) which has an average of two methyl dimethoxy silyl groups at the end was obtained.

(Synthetic example 6)

used, Metallyl end polypropylene oxide was obtained in the same procedure as the synthetic example under the atmosphere of the nitrogen to contain, it was made to react to methyl dimethoxysilane 3.2 group end polypropylene oxide of the number average molecular weight 26,000 [about] produced by polymerizing propylene oxide in the zinc hexa cyanocobaltate glyme complex compound catalyst is Use polyoxypropylene triol of the molecular weight 3.000 [$about\,]$ as an initiator, and the hydroxyl platinum vinyl siloxane complexes isopropanol solution is made into a catalyst to this metallyl end polypropylene oxide 100 weight section, oxygen --- 6vol% --- mixed sulfur at a rate of 1 eq/Pt1eq l except making an allyl chloride into chloridation metallyl. 0.5 copy of platinum content 3wt% of weight section at 90 ** for 5 hours, and the polyoxyalkylene series polymer (A-6) which has an average of 2.8 methyl dimethoxy silyl groups at the end was obtained.

BASA tick 10)) 1.2 weight section and amine (Wako Pure Chemical Industries make, lauryl amine) 0.75 Organic polymer (A-1, A-4-6) 100 weight section which has the reactive silicon group obtained in the Kusumoto Chemicals], DISUPARON 6500) duplexs, and light stabilizer (the Sankyo make.) SANORU SP) 1 weight section, As the amount part of dehydrator vinyltrimetoxysilane (Nippon Unioar make, A-(Japanese east transformation make, neo decanoic acid tin (divalent) (trade name: U-50)) 3.4 weight synthetic example 1 and the synthetic examples 4-6 according to the combination formula shown in 3271 weight section, antioxidant (product made from Ouchi Shinko Chemical industry, NOKURAKKU "S7701 weight section and an ultraviolet ray absorbent (made in Tiba Specialty Chemicals.) Tinuvin weight section are added, After kneading in the state where moisture does not exist substantially Hakuenka CCR) 120 weight section, Titanium oxide (Ishihara Sangyo make, TIPAQUE R-820) 20 section, Carboxylic acid (product made from Japan epoxy resin, neo decanoic acid (trade name: 171) duplexs, adhesion grant agent N-beta-(aminoethyl)-gamma-aminopropyl trimethoxysilane (Nippon Unicar make, A–1120) 3 weight section, and a curing catalyst, Carboxylic acid tin salt under drying conditions, it sealed in the dampproof container and 1 liquid mold-curing nature weight section, DIDP55 weight section, the amount part of thixotropic grant agent (made in Table 3, Surface treatment colloid calcium carbonate (product made from Shiraishi industry, (The reference examples 12-14 and comparative example 6) constituent was obtained.

It examined by having pulled by the same method as the above-mentioned using the class product of Table 3, and was extended at the time of intensity (MPa) and Eb:fracture at the time of M50:50% hauling modulus (MPa) and Tb:fracture, and (%) was measured. A result is shown in Table 3. The recovery was measured by the same method as the above-mentioned using the class product of Table 3. However, the stretched state was fixed at 60 ** 100% for 24 hours this time, and the recovery was measured from the rate which opened this wide at 23 ** and the marked line restored 1 hour afterward. A result is shown in Table 3. The displacement difference of the distance between the marked lines of 200 hours after immediately after performing creep measurement using the piece of a dumbbell, and imposing load using the class

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product of Table 3, by the same method as the method of the reference example 1 and working example 2-4, was measured. A result is shown in Table 3.

Table 3

209	273	483	109	(%)		ΕP		
20.2	78.1	2, 30	5.39	(RPA)		dТ		
0.38	89.0	0.43	0.35	(MPa)		0 9 W	極化物物性	
3.2	1.3	2.2	5.5	(ШШ)		ا ا		
EL	98	T8	87	(%)		率元	歕	
9L '0	97.0	97.0	9 <i>L</i> '0	くきとれりもで		くきて		
2.1	2.1	1.2	7.1	0146744- 3	\I	鑚く、木小は		
3.4	3.4	3.4	3.4	09-N<&Y \$	2	計、XX額、、本小t	東・神小野	
3	3	3	3	A-1120		済 きかま	竹蕎 耔	
7	7	7	7	171-A		隋水	猟	
Į.	ļ	<u> </u>	1	d\$ <i>44€41</i>		除土初	가麵	
Į į	L	1	1	£3£, >357		 原 	條代業	
	ļ	L L	1	0 <i>LL</i> -\$71/-/		廃宝5		
7	7	7	7	ソソ, ロン #6500	} <u>∓</u>	性付与剤	1.4±	
99	99	99	99	DIDP		廃壁	ㅂ	
70	70	70	70	V~. −4K−850	16			
150	120	120	120	ROO華建白		林真	. .	
100				即31	<i>†</i> − ∀			
	100			图8.2	9 – ∀			
		100		2 3個	ĭ – ∀			
			100	图-0.2	3 – ∀	长颓(SA)	 	
9	71	13	15	基素トヤガふ灵				
脉棘出		- 例字零	A	ወሀታ ል ችሉ!		(陪量重) 海眛		

that the organic polymer with many reactive silicon groups per molecule (A-1, A-5-6) is excellent in Comparison with the reference examples 12-14 of Table 3 and the comparative example 6 shows stability and creep resistance.

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(Synthetic example 7)

used, Metallyl end polypropylene oxide was obtained in the same procedure as the synthetic example 6. To this metallyl end polypropylene oxide, in the same procedure as the synthetic example 6, it was group end polypropylene oxide of the number average molecular weight 28,500 [about] produced by made to react to methyl dimethoxysilane and the polyoxyalkylene series polymer (A-7) which has an Use polyoxypropylene glycol of the molecular weight 2,000 [about] as an initiator, and the hydroxyl polymerizing propylene oxide in the zinc hexa cyanocobaltate glyme complex compound catalyst is average of 1.9 methyl dimethoxy silyl groups at the end was obtained.

(Synthetic example 8)

To the metallyl end polypropylene oxide obtained in the synthetic example 7, in the same procedure as the synthetic example 6, it was made to react to methyl dimethoxysilane and the polyoxyalkylene series polymer (A-8) which has an average of 1.5 methyl dimethoxy silyl groups at the end was obtained.

(Synthetic example 9)

To this allyl end polypropylene oxide, in the same procedure as the synthetic example 1, it was made group end polypropylene oxide of the number average molecular weight 28,500 [about] produced by Use polyoxypropylene glycol of the molecular weight 2,000 [about] as an initiator, and the hydroxyl used, Allyl end polypropylene oxide was obtained in the same procedure as the synthetic example 1. polymerizing propylene oxide in the zinc hexa cyanocobaltate glyme complex compound catalyst is to react to methyl dimethoxysilane and the polyoxyalkylene series polymer (A-9) which has an average of 1.5 methyl dimethoxy silyl groups at the end was obtained.

(The <u>reference examples 15-16</u> and comparative examples 7-8) Organic polymer (A-4, A-7-9) 100 weight section which has the reactive silicon group obtained in the weight section, DIDP55 weight section, the amount part of thixotropic grant agent (made in [Kusumoto Chemicals], DISUPARON 6500) duplexs, and light stabilizer (the Sankyo make.) SANORU synthetic example 4 and the synthetic examples 7-9 according to the combination formula shown in Hakuenka CCR) 120 weight section, Titanium oxide (Ishihara Sangyo make, TIPAQUE R-820) 20 Table 4, Surface treatment colloid calcium carbonate (product made from Shiraishi industry,

dibutyitin bisacetylacetonate (Japanese east transformation make, neo SUTAN U-220) duplexs were 3271 weight section, antioxidant (product made from Ouchi Shinko Chemical industry, NOKURAKKU -S7701 weight section and an ultraviolet ray absorbent (made in Tha Specialty Chemicals.) Tinuvin 171) duplexs, adhesion grant agent N-beta-(aminoethyl)-gamma-aminopropyl trimethoxysilane (the SP) 1 weight section, The amount part of dehydrator vinyltrimetoxysilane (Nippon Unicar make, Aconditions, it sealed in the dampproof container and 1 liquid mold-curing nature constituent was Nippon Unicar make.) A-1120) Three weight sections and the amount part of curing catalyst added, after kneading in the state where moisture does not exist substantially under drying

It examined by having pulled by the same method as the above-mentioned using the class product of Table 4, and was extended at the time of intensity (MPa) and Eb:fracture at the time of M50:50% hauling modulus (MPa) and Tb:fracture, and (%) was measured. A result is shown in Table 4.

The recovery was measured by the same method as the above-mentioned using the class product of recovery was measured from the rate which opened this wide at 23 ** and the marked line restored Table 4. However, the stretched state was fixed at 23 ** 100% for 24 hours this time, and the 24 hours afterward. A result is shown in Table 4.

after performing creep measurement using the piece of a dumbbell, and imposing load using the class product of Table 4, by the same method as the method of the reference example 1 and working example 2-4, was measured. A result is shown in Table 4.

The displacement difference of the distance between the marked lines of 45 hours after immediately

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[Table 4]

403	8101	6101	652		(%)		qョ		
1.90	2.60	3, 02	2, 71	(RPA)		ЧT			
65.0	0.19	0.21	0.41		(69M)		0 g W	确化物物体	
17	44	77	GL		(ww)		4-1		
G/	ÞĹ	1/8	98		(%)		率元		
7	7	7	7		02 2-U <&X X \$		欺 嫡:		
3	3	3	ε		0S11-A		 		
7	7	7	7		171-A		1		
Ţ	Ĺ	L	Ţ		d\$4% <u>≤</u> 47		[降117]		
l '	l	1	Į.		下25ペ・ン327				
Ţ	ļ	I	Ţ	OFF-2.M-\# N N 宝光					
7	7	7	7	0	099# ベロ ゚ハスト ゙テ	降も付置して			
99	99	99	99		9010				
20	50	50	50		<i>4√</i> ~.− <i>4B</i> −850]		
150	150	150	150		ROO華體白			茶	
100				剧 3 1	基机气	<i>†</i> – ∀			
	100			里·S·L	基化亿	6 – ∀]	
""		001		1 2個	基机铁	8 – A]		
			001	凰6 1	章40.6%	$\nabla - A$	长孰(£A)	朴合重勬育	
8	L	91	91	基素下寸對初凤	煮耕の基齢末の 値			_	
(科 交	車和	[4]	¥ ≨	ወፀታልችሉ!	人草基素內對卻反		(陪量重)		

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Comparison with the reference examples 15~16 of Table 4 and the comparative examples 7~8 shows that the organic polymer (A-7-8) which introduced the reactive silicon group to the metallyl group end organicity polymer is excellent in stability and creep resistance.

(Synthetic example 10)

To the allyl end polypropylene oxide obtained in the synthetic example 1, in the same procedure as the synthetic example 1, it was made to react to triethoxysilane and the polyoxyalkylene series polymer (A-10) which has an average of 2.3 triethoxy silyl groups at the end was obtained.

(The reference example 17 and the comparative examples 9-10)

made from Ouchi Shinko Chemical industry, NOKURAKKU SP) 1 weight section, As a dehydrator, the Organic polymer (A-1, A-10) 100 weight section which has the reactive silicon group obtained in the (Japanese east transformation make, neo SUTAN U-220) duplexs, After kneading in the state where Table 5, Surface treatment colloid calcium carbonate (product made from Solvay, Winnoffl SPM) 120 weight section, Titanium oxide (product made from Kerr-McGee, RFK-2) 20 weight section, DIUP50 weight section, Thixotropic grant agent (product made from Cray Valley, Crayvallac super) 5 weight triethoxysilane (the Nippon Unicar make.) which is the (G) ingredient as an adhesion grant agent Amoisture does not exist substantially under drying conditions, it sealed in the dampproof container synthetic example 1 and the synthetic example 10 according to the combination formula shown in 1100) Or add N-beta-(aminoethyl)-gamma-aminopropyl trimethoxysilane (Nippon Unicar make, Aabsorbent (made in Tiba Specialty Chemicals.) Tinuvin 3271 weight section, antioxidant (product amount part of vinyltrimetoxysilane (Nippon Unicar make, A-171) duplexs, gamma-aminopropyl section, light stabilizer (Sankyo make, SANORULS770) 1 weight section and an ultraviolet ray 1120) 3 weight section and the amount part of ouring catalyst dibutyitin bisacetylacetonate and 1 liquid mold-curing nature constituent was obtained. The recovery was measured by the same method as the above-mentioned using the class product of recovery was measured from the rate which opened this wide at 23 ** and the marked line restored Table 5. However, the stretched state was fixed at 60 ** 100% for 24 hours this time, and the I hour afterward, A result is shown in Table 5.

the method of the <u>reference examples</u> 5-11, using the class product of Table 5, was measured. As for The displacement difference of the distance between the marked lines of 140 hours after immediately after performing creep measurement using a shear sample and imposing load by the same method as the valuation basis, the displacement difference made O x for the thing below 0.4 mm, and the displacement difference carried out a thing of 0.4 mm or more. A result is shown in Table 5.

(Hardenability of a hardenability constituent)

covered time) until the surface stretches a hide under 23 ** and 50% of humidity RH conditions was The class product of Table 5 was thinly lengthened in thickness of about 3 mm, and time (leathermeasured. The one where leather-covered time is shorter means that hardenability is excellent. A result is shown in Table 5.

[Table 5]

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if the aminosilane which has a triethoxy silyl group which is the (G) ingredient as an adhesive grant

agent is combined with the er group as an organic polymer and creep resistance, change good. [0356]	(The <u>reference example</u> 18 ar Organic polymer (A-2) 100 w synthetic example 2 accordin section and a dehydrator — t aminopropyl trimethoxysilane	
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and of the (A4) ingredient using the polymer which has a triethoxy silyl as shown in the reference example 17 of Table 5, Excelling in stability e of the skinning time in storage order is small, and storage stability is

nitrogen purge, and 1 liquid mold-curing nature constituent was obtained. In the comparative example transformation make, neo SUTAN U-220) duplexs were sealed in the glassware which carried out the 11, the leather-covered time test was performed under 50% of 23 ** humidity RH conditions, without recuperating oneself in this 1 liquid mold-curing nature constituent. In the <u>reference example</u> 18 and the comparative example 12, after promoting the ester exchange reaction between reactive silicon triethoxysilane (made in a col coat.) the N-beta-(aminoethyl)-gammae (the Nippon Unicar make.) which is the (H) ingredient as the amount gamma-aminopropyl triethoxysilane (the Shin-Etsu Chemical make.) It added, KBE-6033 weight ng to the combination formula shown in Table 6, as DIDP30 weight exs, and an adhesion grant agent A-1120) or N-beta-(aminoethyl)weight section which has the reactive silicon group obtained in the section and the amount part of curing catalyst dibutyltin bisacetylacetonate (Japanese east groups by recuperating oneself for seven days at 50 ** in these 1 liquid mold-curing nature constituents, the leather-covered time test was performed under 50% of 23 ** humidity RH and the comparative examples 11–12) conditions. A result is shown in Table 6.

Table 6

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used for the end of the (A4) ingredient as an organic polymer, If the aminosilane which has a methoxy sily group which is the (H) ingredient as an adhesive grant agent is combined and an ester exchange reaction is promoted by care of health, the hardenability of an organic polymer can be raised notably. As shown in the reference example 18 of Table 6, the polymer which has a triethoxy silyl group is

(The reference examples 19~20 and comparative example 13)

ndustry, Hakuenka CCR) 120 weight section. Titanium oxide (Ishihara Sangyo make, TIPAQUE R-820) [Kusumoto Chemicals], DISUPARON 6500) duplexs, and light stabilizer (the Sankyo make.) SANORU catalyst of the (E) ingredient as a curing catalyst what carried out concomitant use addition of BASA bisacetylacetonate (Japanese east transformation make, neo SUTAN U-220) duplexs was made into 3271 weight section, antioxidant (product made from Ouchi Shinko Chemical industry, NOKURAKKU LS7701 weight section and an ultraviolet ray absorbent (made in Tiba Specialty Chemicals.) Tinuvin catalysts were added, after kneading in the state where moisture does not exist substantially under 171) duplexs, adhesion grant agent N-beta-(aminoethyl)-gamma-aminopropyl trimethoxysilane (the was obtained, the neo decanoic acid (the product made from Japan epoxy resin.) which is a non-tin SP) 1 weight section, The amount part of dehydrator vinyltrimetoxysilane (Nippon Unicar make, Adrying conditions, it sealed in the dampproof container and 1 liquid mold-curing nature constituent Matsumoto Trading make.) What carried out Olga Chicks TC-750 8.5 weight-section addition was weight section — the reference example 19 and isopropoxy titanium bis (ethylacetoacetate) (the synthetic example 10, Surface treatment colloid calcium carbonate (product made from Shiraishi Organic polymer (A-10) 100 weight section which has the reactive silicon group obtained in the tick 106 weight section and the amine (Wako Pure Chemical Industries make, lauryl amine) 0.75 20 weight section, DIDP55 weight section, the amount part of thixotropic grant agent (made in Nippon Unicar make,) A-1120) Three weight sections and the various below-mentioned curing made into the reference example 20. What carried out the amount part addition of dibutyltin the comparative example 13.

class products, the hardened material of the <u>reference example 19</u> and the <u>reference example 20</u> showed the recovery higher than the hardened material of the comparative example 13. As a result of measuring the recovery by the same method as the above-mentioned using these

(Synthetic example 11)

to react to triethoxysilane and the polyoxyalkylene series polymer (A-11) which has an average of 1.5 group end polypropylene oxide of the number average molecular weight 25,500 [about] produced by To this allyl end polypropylene oxide, in the same procedure as the synthetic example 1, it was made Use polyoxypropylene glycol of the molecular weight 2,000 [about] as an initiator, and the hydroxyl used, Allyl end polypropylene oxide was obtained in the same procedure as the synthetic example 1. polymerizing propylene oxide in the zinc hexa cyanocobaltate glyme complex compound catalyst is triethoxy silyl groups at the end was obtained.

(Synthetic example 12)

To the allyl end polypropylene oxide obtained in the synthetic example 11, in the same procedure as the synthetic example 1, it was made to react to methyl dimethoxysilane and the polyoxyalkylene series polymer (A-12) which has an average of 1.5 methyl dimethoxy silyl groups at the end was obtained.

(The reference example 21 and the comparative examples 14-15)

section, thixotropic grant agent (made in [Kusumoto Chemicals], DISUPARON 305) 3 weight section, calcium-carbonate (product made from Shiroishi calcium, HOWAITON SB) 20 weight section, DOP40 calcium carbonate (product made from Shiraishi industry, BISUKO light R) 60 weight section, Heavy-Organic polymer (A-11, A-12) 95 weight section which has the reactive silicon group obtained in the weight section, epoxy system plasticizer (New Japan Chemical make, SANSO sizer EP-S) 20 weight synthetic example 11 and the synthetic example 12, Surface treatment colloid calcium carbonate (product made from Shiraishi industry, Hakuenka CCR) 60 weight section, Surface treatment colloid

and a photo-setting resin (the Toagosei make.) ARONIKKUSU M-3093 weight section, light stabilizer (Sankyo make, SANORULS770) 1 weight section, Ultraviolet ray absorbent (made in [Tiba Specialty Chemicals], tinuvin 327) 1 weight section, Zero copy of minute hollow body (the product made from the Fuji SHIRISHIA chemicals, the FUJIBA lune H-40) which is antioxidant (made in [Tiba Specialty Chemicals], IRUGA NOx 1010) 1 weight section and the (F) ingredient, or 20 copies were measured, respectively, and it often kneaded with a 3 paint roll, and was considered as base resin. What added 20 copies of minute hollow bodies was made into the comparative example 12, using (A-12) as an organic polymer, and what added 20 copies of minute hollow bodies was made into the comparative example 15, using (A-11) as an organic polymer.

Using the mixture of 2-ethylhexanoic acid tin (Japanese east transformation make, U-28) (divalent) 3 weight section and amine (Wako Pure Chemical Industries make, lauryl amine) 0.75 weight section as a hardening agent, above-mentioned base resin and hardening agent were mixed uniformly, and workability (******** and endurance were evaluated.

The constituent of the <u>reference example</u> 21 had workability better than the comparative example 14, and its endurance was better than the comparative example 15.

(The reference example 22 and the comparative example 16)

carbonate (product made from Shiraishi industry, BISUKO light R) 60 weight section, Heavy-calcium-carbonate (product made from Shiroishi calcium, HOWAITON SB) 20 weight section, DOP40 weight section, epoxy system plasticizer (New Japan Chemical make, SANSO sizer EP-S) 20 weight section, RUGA NOx 1010) I weight section were measured, respectively, and it often kneaded with a 3 paint example 10, or the synthetic example 1, Surface treatment colloid calcium carbonate (product made thixotropic grant agent (made in [Kusumoto Chemicals], DISUPARON 305) 3 weight section, and a (Sankyo make, SANORULS770) 1 weight section, Ultraviolet ray absorbent (made in [Tiba Specialty made into the <u>reference example 22,</u> and what added 95 copies of (A-1) as an organic polymer was roll, and was considered as base resin. What added 70 copies of (A-10) as an organic polymer was amine (Wako Pure Chemical Industries make, lauryl amine) 0.75 weight section was added, and the made into the comparative example 16.As a hardening agent, to this base resin, the mixture of 2ethylhexanoic acid tin (Japanese east transformation make, U-28) (divalent) 3 weight section and Organic polymer (A-1) 95 weight section which has the reactive silicon group obtained in organic Chemicals], tinuvin 327) 1 weight section and antioxidant (made in [Tiba Specialty Chemicals], polymer (A-10) 70 weight section which has the reactive silicon group obtained in the synthetic photo-setting resin (the Toagosei make.) ARONIKKUSU M-3093 weight section, light stabilizer from Shiraishi industry, Hakuenka CCR) 60 weight section, Surface treatment colloid calcium recovery was measured to it.

The constituent of the <u>reference example</u> 22 showed the recovery higher than the comparative comparative seample 16, stopping weight % of an organic polymer low.

(The reference example 23 and the comparative example 17)

Organic polymer (A-10) 95 weight section which has the reactive silicon group obtained in the synthetic example 10, Surface treatment colloid calcium carbonate (product made from Shiraishi industry, Hakuenka CCR) 60 weight section, Surface treatment colloid calcium carbonate (product made from Shiraishi industry, BISUKO light R) 60 weight section, Heavy-calcium-carbonate (product made from Shiroishi calcium, HOWAITON SB) 20 weight section, DOP40 weight section, epoxy system plasticizer (New Japan Chemical B, DISUPARON 305) 3 weight section, thixotropic setting resin (the Toagosei make, ANSO sizer EP-S) 20 weight section, thixotropic setting resin (the Toagosei make, ANSO WIKKUSU M-3093 weight section, light stabilizer (Sankyo make, SANORULS770) 1 weight section, Ultraviolet ray absorbent (made in [Tiba Specialty Chemicals], tinuvin 327) 1 weight section, Antioxidant (made in [Tiba Specialty Chemicals], tinuvin 327) 1 weight section, Antioxidant (made in [Tiba Specialty Chemicals], IRUGA NOX 1010) 1 weight section and zero copy of epoxy resin (the product made from Japan epoxy resin, Epicoat 828), or five copies were measured, respectively, and it often kneaded with a 3 paint roll, and

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was considered as base resin. What added five copies of epoxy resins was made into the reference example 17. As a hardening agent, to this base resin, the mixture of 2-ethylhexanoic acid tin (Japanese east transformation make, U-28) (divalent) 3 weight section and amine (Wako Pure Chemical Industries make, lauryl amine) 0.75 weight section was added, and the recovery was measured to it.

The constituent of the <u>reference example</u> 23 showed the recovery higher than the comparative

example 17.

The reference example 24 and the comparative example 18)

Organic polymer (A–10) 95 weight section which has the reactive silioon group obtained in the synthetic example 10. Surface treatment colloid caloium carbonate (product made from Shiraishi industry, Hakuenka CCR) 60 weight section, Surface treatment colloid caloium carbonate (product made from Shiraishi industry, BISUKO light R) 60 weight section, Heavy-caloium-carbonate (product made from Shiraishi industry, BISUKO light R) 60 weight section, Heavy-caloium-carbonate (product made from Shiraishi caloium, HOWAITON SB) 20 weight section, DOP40 weight section, epoxy system plasticizer (New Japan Chemical B) DISUPARON 305) 3 weight section, thixotropic grant agent (made in [Kusumoto Chemicals], DISUPARON 305) 3 weight section, thixotropic setting resin (the Toagosei make) ARONIKKUSU M-3093 weight section, light stabilizer (Sankyo make, SANORULS770) 1 weight section, Ultraviolet ray absorbent (made in [Tiba Specialty Chemicals], Ghemicals], Itinuvin 327) 1 weight section and antioxidant (made in [Tiba Specialty Chemicals], IRUGA NOx 1010) 1 weight section and antioxidant (made in [Tiba Specialty Spaint roll, and was considered as base resin.

Pethylhexanoic acid tin (Japanese east transformation make, U-28) (divalent) 3 weight section and amine (the Wako Pure Chemical Industries make.) lauryl amine 0.75 weight section and dibutyltin bisacetylacetonate (the Japanese east — transformation — make.) The thing using the mixture of neo SUTAN U-220 0.1 weight section as a hardening agent is made into the reference example 24. The thing using the mixture of 2-ethylhexanoic acid tin (Japanese east transformation make, U-28) (divalent) 3 weight section and amine (Wako Pure Chemical Industries make, lauryl amine) 0.75 weight section as a hardening agent was made into the comparative example 18. Base resin and a hardening agent, were mixed uniformly and the recovery and thin layer hardenability were evaluated.

The constituent of the reference example 24 showed good thin layer hardenability rather than the comparative example 18, while the high recovery was shown.

[0373]

(Synthetic example 13)

To the allyl end polyisobutylene obtained according to the example of manufacture of JP,H11–209639,A, under existence of Pt catalyst, it was made to react to triethoxysilane and the polyisobutylene (A–13) which has a triethoxy silyl group at the end was obtained.

[0374]

(Synthetic example 14)

To the allyl end polyisobutylene obtained in the synthetic example 13, under existence of Pt catalyst, it was made to react to methyl dimethoxysilane and the polyisobutylene (A-14) which has a methyl dimethoxy silyl group at the end was obtained.

(The reference example 25 and the comparative example 19)

To organic polymer (A-13, A-14) 100 weight section which has the reactive silicon group obtained in the synthetic example 13 and the synthetic example 14, the amount part of dibuty/tin bisacetylacetonate (Japanese east transformation make, neo SUTAN U-220) duplexs was added, and the hardened material was obtained. The thing using (A-13) as an organic polymer was made into the reference example 25, and the thing using (A-14) was made into the comparative example 19. The hardened material of the reference example 25 showed the recovery higher than the comparative example 19.

[0376]

(Synthetic example 15)

Pentamethyl diethylenetriamine (0.17g) was added and the polymerization was made to start. Acrylic CuBr (4.2g) and acetonitrile (27.3g) were added to the reaction vessel with an agitator, and it stirred acid n-butyl (400g) was dropped continuously, heating and stirring at 70 **. Dividing addition of the diethyl adipate (8.8g), and acetonitrile (16.6g) were added to this, and stirring mixing was improved. for 15 minutes at 65 ** under a nitrogen atmosphere. Acrylic acid n-butyl (100g), 2, 5-dibromo triamine (0.68g) was carried out in the middle of dropping of acrylic acid n-butyl.

10 ** succeedingly, and the mixture containing the polymer which has an alkenyl group was obtained **, 1,7-octadien (53.7g), acetonitrile (132g), and triamine (1.69g) were added, it heated and stirred at When monomer conversion reaches to 96%, after devolatilizing a ** monomer and acetonitrile at 80

added to the methyloyclohexane solution of the polymer to 100 copies of polymers, and it heated and copies of KYO word 700SL [:] product [Both] made from Harmony Chemicals) of adsorbent was deating devolatilization of acetonitrile in a mixture and the unreacted 1,7-octadien was carried out, polymer (polymer [P1]) which has an alkenyl group by condensing a polymer solution was obtained. sediment with a centrifuge, and was removed. Six copies (three copies of KYO word 500SH $^\prime$ 3 stirred under oxygen and nitrogen mixed gas atmosphere. Insoluble matter was removed and the and it diluted with the methylcyclohexane. The insoluble polymerization catalyst was made to

100 copies of the polymer with 400 copies of methyloyclohexanes further and removing solid content, After having carried out heating devolatilization (10 or less torr of decompression degrees), diluting polymer [P2] was obtained. The number average molecular weight of this polymer [P2] was 24800, and molecular weight distribution was 1.36. The number of the alkenyl groups introduced per one stirring the obtained polymer [P1] at 180 ** for 12 hours, the solution was condensed and the molecule of polymers was 1.8.

were added in order, and it mixed, and heated and stirred at 100 ** under a nitrogen atmosphere for ethynyl)-1, 1 and 3, and 3-tetramethyl disiloxane (they are 1.5 mol equivalents to an alkenyl group) trimethoxysilyl group content polymer (A-15) which condenses a reaction mixture and is made into distribution was 1.32. The number of the silyl groups introduced per one molecule of polymers was the purpose was obtained. The number average molecular weight was 27900 and molecular weight To this polymer [P2], methyl orthoformate (it is 1 mol equivalent to an alkenyl group), A platinum 0.5 hour. It checked that the alkenyl group had disappeared by a reaction by ¹H-NMR, and the catalyst (it is 10 mg to 1 kg of polymers as an amount of platinum metal), 1-(2-trimethoxysilyl

(Synthetic example 16)

average molecular weight was 28600 and molecular weight distribution was 1.48. The number of the ethynyl)–1 used in the synthetic example 15, 1 and 3, and 3-tetramethyl disiloxane. The number As opposed to the polymer [P2] obtained in the synthetic example 15, The triethoxy silyl group triethoxysilane (they are 3 mol equivalents to an alkenyl group) instead of 1-(2-trimethoxysilyl content polymer (A-16) was obtained like the synthetic example 15 except having used silyl groups introduced per one molecule of polymers was 1.5.

(Synthetic example 17)

an alkenyl group,) instead of 1-(2-trimethoxysilyl ethynyl)-1 used in the synthetic example 15, 1 average molecular weight was 28400 and molecular weight distribution was 1.51. The number of the As opposed to the polymer [P2] obtained in the synthetic example 15. It is methyl dimethoxysilane (to an alkenyl group.) instead of 1-(2-trimethoxysilyl ethynyl)-1 used in the synthetic example 15. I and 3, and 3-tetramethyl disiloxane. The methyl dimethoxy silyl group content polymer (A-17) was obtained like the synthetic example 15 except having used three mol equivalents. The number silyl groups introduced per one molecule of polymers was 1.5.

as opposed to organic polymer 100 weight section which has a reactive silicon group -- surface treatment colloid calcium carbonate (the product made from the Shiraishi industry.) Hakuenka (The reference examples 26-28 and comparative example 20)

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(A-4) 50 weight section which has the methyl dimethoxy silyl group obtained in the synthetic example the dampproof container and 1 liquid mold-curing nature constituent was obtained. The **** thing for kneading in the state where moisture does not exist substantially under drying conditions, it sealed in which has a reactive silicon group, (A-15) Make the *** thing for a total of 100 weight sections into 4, The **** thing for 100 weight sections is made into the reference example 28 for the acrylio ester system polymer (A-17) which has the methyl dimethoxy silyl group obtained in the synthetic example 20 weight section, titanium oxide (Ishihara Sangyo make, TIPAQUE R-820) 10 weight section, DIDP60 system polymer (A-16) which has the triethoxy silyl group obtained in the synthetic example 16, The 100 weight sections is made into the reference example 26 for the acrylic ester system polymer (A-15) which has the trimethoxysilyl group obtained in the synthetic example 15 as an organic polymer DISUPARON 6500) duplexs, light stabilizer (Sankyo make, SANORULS765) 1 weight section and an adhesion grant agent N-beta-(aminoethyl)-gamma-aminopropyl trimethoxysilane (the Nippon Unicar make.) A–1120) Dibutyltin bisacetylaoetonate (Japanese east transformation make, neo SUTAN Uthe reference example 27 for the mixture of 50 weight sections and polyoxyalkylene series polymer **** thing for 100 weight sections was made into the comparative example 20 for the acrylic ester CCR150 weight section and heavy calcium carbonate (the Maruo Calcium make.) 25ANANOKKUSU dehydrator vinyltrimetoxysilane (the Nippon Unicar make.) A-171) the amount part of duplexs, and 17. The hardened material of the reference examples 26-28 showed the recovery higher than the 220) 0.2 weight section was added as the amount part of duplexs, and a curing catalyst, after ultraviolet ray absorbent (made in Tiba Specialty Chemicals.) tinuvin 2131 weight section and weight section, the amount part of thixotropic grant agent (made in [Kusumoto Chemicals], comparative example 20.

Effect of the Invention]

The hardenability constituent of this invention is excellent in stability, endurance, and creep

[Translation done.]